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- (57) Abstract

The present invention provides a fusion polypeptide capable of binding a cytokine to form a nonfunctional complex. It also provides a nucleic acid sequence encoding the fusion polypeptide and methods of making and uses for the fusion polypeptide.

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RECEPTOR BASED ANTAGONISTS AND METHODS OF MAKING AND USING

This application claims priority of U.S. Application No. 09/313,942, filed May 19, 1999, which claims priority of U.S. Provisional Application No. 60/101,858 filed September 25, 1998. Throughout this application various publications are referenced. The disclosures of these publications in their entireties are hereby incorporated by reference into this application.

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BACKGROUND OF THE INVENTION

Although discovered for varying biological activities, ciliary neurotrophic factor (CNTF), leukemia inhibitory factor (LIF), oncostatin M (OSM) and interleukin-6 (IL-6) comprise a defined family of cytokines (referred to herein as the "CNTF family" of cytokines). These cytokines are grouped together because of their distant structural similarities [Bazan, J. Neuron 7: 197-208 (1991); Rose and Bruce, Proc. Natl. Acad. Sci. USA 88: 8641-8645 (1991)], and, perhaps more importantly, because they share "β" signaltransducing receptor components [Baumann, et al., J. Biol. Chem. 265:19853-19862 (1993); Davis, et al., Science 260: 1805-1808 (1993); Gearing et al., Science 255:1434-1437 (1992); Ip et al., Cell 69: 1121-1132 (1992); Stahl, et al., J. Biol. Chem. 268: 7628-7631 (1993); Stahl and Yancopoulos, Cell 74: 587-590 (1993)]. Receptor activation by this family of cytokines results from either homo- or hetero-dimerization of these β components [Davis, et al. Science 260: 1805-1808 (1993), Murakami, et al., Science 260: 1808-1810 (1993); Stahl and Yancopoulos, Cell 74: 587-590 (1993)]. IL-6 receptor activation requires homodimerization of gp130 [Murakami, et al. Science 260: 1808-1810 (1993), Hibi, et al., Cell 63: 1149-1157 (1990)], a protein initially identified as the IL-6 signal transducer [Hibi, et al., Cell 63: 1149-1157 (1990)]. CNTF, LIF and OSM receptor activation results from heterodimerization between gp130 and a second gp130-related protein known as LIFR\$ [Davis,

et al., Science 260: 1805-1808 (1993)], that was initially identified by its ability to bind LIF [Gearing et al., EMBO J. 10: 2839-2848 (1991)].

In addition to the β components, some of these cytokines also require specificity-determining " α " components that are more limited in their tissue distribution than the β components, and thus determine the cellular targets of the particular cytokines [Stahl and Yancopoulos, Cell 74: 587-590 (1993)]. Thus, LIF and OSM are broadly acting factors that may only require the presence of gp130 and LIFR β on responding cells, while CNTF requires CNTFR α [Stahl and Yancopoulos, Cell 74: 587-590 (1993)] and IL-6 requires IL-6R α [Kishimoto, et al., Science 258: 593-597 (1992)]. Both CNTFR α (Davis et al., Science 259:1736-1739 (1993) and IL-6R α [Hibi, et al. Cell 63:1149-1157, Murakami, et al., Science 260:1808-1810 (1990); Taga, et al., Cell 58:573-581 (1989)] can function as soluble proteins, consistent with the notion that they do not interact with intracellular signaling molecules but that they serve to help their ligands interact with the appropriate signal transducing β subunits [Stahl and Yancopoulos, Cell 74: 587-590 (1993)].

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Additional evidence from other cytokine systems also supports the notion that dimerization provides a common mechanism by which all cytokine receptors initiate signal transduction. Growth hormone (GH) serves as perhaps the best example in this regard. Crystallographic studies have revealed that each GH molecule contains two distinct receptor binding sites, both of which are recognized by the same binding domain in the receptor, allowing a single molecule of GH to engage two receptor molecules [de Vos, et al., Science 255: 306-312 (1992)]. Dimerization occurs sequentially, with site 1 on the GH first binding to one receptor molecule, followed by the binding of site 2 to a second receptor molecule [Fuh, et al., Science 256: 1677-1680 (1992)]. Studies with the erythropoietin (EPO) receptor are also consistent with the importance of dimerization in receptor activation, as EPO receptors can be constitutively activated by a

single amino acid change that introduces a cysteine residue and results in disulfide-linked homodimers [Watowich, et al., Proc. Natl. Acad. Sci. USA 89:2140-2144 (1992)].

In addition to homo- or hetero-dimerization of β subunits as the critical 5 step for receptor activation, a second important feature is that formation of the final receptor complex by the CNTF family of cytokines occurs through a mechanism whereby the ligand successively binds to receptor components in an ordered manner [Davis, et al. Science 260:1805-1818 (1993); Stahl and Yancopoulos, Cell 74: 587-590 (1993)]. Thus CNTF first 10 binds to CNTFRa, forming a complex which then binds gp130 to form an intermediate (called here the abl intermediate) that is not signaling competent because it has only a single \beta component, before finally recruiting LIFR β to form a heterodimer of β components which then initiates signal transduction. Although a similar intermediate containing 15 IL-6 bound to IL-6Rα and a single molecule of gp130 has not been directly isolated, we have postulated that it does exist by analogy to its distant relative, CNTF, as well as the fact that the final active IL-6 receptor complex recruits two gp130 monomers. Altogether, these findings led to a proposal for the structure of a generic cytokine receptor complex (Figure 1) 20 in which each cytokine can have up to 3 receptor binding sites: a site that binds to an optional α specificity-determining component (α site), a site that binds to the first β signal-transducing component (β 1 site), and a site that binds to the second β signal-transducing component (β 2 site) [Stahl and Yancopoulos, Cell 74: 587-590 (1993)]. These 3 sites are used in 25 sequential fashion, with the last step in complex formation -- resulting in β component dimerization -- critical for initiating signal transduction [Davis, et al. Science 260:1805-1818 (1993)]. Knowledge of the details of receptor activation and the existence of the non-functional β1 intermediate for CNTF has led to the finding that CNTF is a high affinity 30

antagonist for IL-6 under certain circumstances, and provides the strategic basis for designing ligand or receptor-based antagonists for the CNTF family of cytokines as detailed below.

Once cytokine binding induces receptor complex formation, the 5 dimerization of β components activates intracellular tyrosine kinase activity that results in phosphorylation of a wide variety of substrates [Ip, et al. Cell 69:121-1132 (1992)]. This activation of tyrosine kinase appears to be critical for downstream events since inhibitors that block the tyrosine phosphorylations also prevent later events such as gene inductions [Ip, et 10 al., Cell 69:121-1132 (1992); Nakajima and Wall, Mol. Cell. Biol. 11:1409-1418 (1991)]. Recently, we have demonstrated that a newly discovered family of non-receptor tyrosine kinases that includes Jak1, Jak2, and Tyk2 (referred to as the Jak/Tyk kinases) [Firmbach-Kraft, et al., Oncogene 5:1329-1336 (1990); Wilks, et al., Mol. Cell. Biol. 11: 2057-2065 (1991) and that 15 are involved in signal transduction with other cytokines [Argetsinger, et al., Cell 74:237-244 (1993); Silvennoinen, et al., Proc. Natl. Acad. Sci. USA 90:8429-8433 (1993); Velazquez, et al., Cell 70: 313-322 (1992); Witthuhn, et al., Cell 74:227-236 (1993)], preassociate with the cytoplasmic domains of the β subunits gp130 and LIFRβ in the absence of ligand, and become tyrosine 20 phosphorylated and activated upon ligand addition [Stahl et al., Science 263:92-95 (1994)]. Therefore these kinases appear to be the most proximal step of intracellular signal transduction activated inside the cell as a result of ligand binding outside of the cell. Assay systems for screening collections of small molecules for specific agonist or antagonist activities 25 based on this system are described below.

The CNTF family of cytokines play important roles in a wide variety of physiological processes that provide potential therapeutic applications for both antagonists and agonists.

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SUMMARY OF THE INVENTION

An object of the present invention is the production of cytokine antagonists that are useful in the treatment of cytokine-related diseases or disorders.

Another object of the invention is the use of the disclosed cytokine antagonists for the treatment of cytokine-related diseases or disorders. For example, an IL-6 antagonist described herein may be used for the treatment of osteoporosis, the primary and second effects of cancers, including multiple myeloma, or cachexia.

Another object of the invention is the development of screening systems useful for identifying novel agonists and antagonists of cytokine receptors.

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Another object of the invention is the development of screening systems useful for identifying small molecules that act as agonists or antagonists of the cytokines.

Another object of the invention is the development of screening systems useful for identifying novel agonists and antagonists of members of the CNTF family of cytokines.

Another object of the invention is the development of screening systems
useful for identifying small molecules that act as agonists or antagonists of
the CNTF family of cytokines.

BRIEF DESCRIPTION OF THE FIGURES

FIGURE 1: Ordered binding of receptor components in a model of a generic cytokine receptor. The model indicates that cytokines contain up to 3 receptor binding sites and interact with their receptor components by

binding first the optional α component, followed by binding to $\beta 1$, and then $\beta 2$. The β components for many cytokine receptors interact through membrane proximal regions (shaded boxes) with the Jak/Tyk family of cytoplasmic protein tyrosine kinases. Only upon dimerization of β components is signal transduction initiated, as schematized by the tyrosine phosphorylations (P) of the β components and the Jak/Tyk kinases.

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- FIGURE 2: CNTF inhibits IL-6 responses in a PC12 cell line (called PC12D) that expresses IL6Rα, gp130, CNTFRα, but not LIFRβ. Serum-deprived PC12D cells were incubated + IL-6 (50 ng/mL) in the presence or absence of CNTF as indicated. Some plates also received soluble IL6Rα (1 mg/mL) or soluble CNTFRα (1 mg/mL) as indicated. Cell lysates were subjected to immunoprecipitation with anti-gp130 and immunoblotted with anti-phosphotyrosine. Tyrosine phosphorylation of gp130 is indicative of IL-6 induced activation of the IL-6 receptor system, which is blocked upon coaddition of CNTF.
- FIGURE 3: Scatchard analysis of iodinated CNTF binding on PC12D cells. PC12D cells were incubated with various concentrations of iodinated CNTF in the presence or absence of excess non-radioactive competitor to determine the specific binding. The figure shows a Scatchard plot of the amount of iodinated CNTF specifically bound, and gives data consistent with two binding sites with dissociation constants of 9 pM and 3.4 nM.
- FIGURE 4. The amino acid sequence of human gp130-Fc-His6. Amino acids 1 to 619 are from human gp130 (Hibi et al., Cell 63:1149-1157 (1990). Note that amino acid number 2 has been changed from a Leu to a Val in order to accommodate a Kozak sequence in the coding DNA sequence. The signal peptide of gp130-Fc-His6 has been italicized (amino acids 1 to 22). The Ser-Gly bridge is shown in bold type (amino acids 620, 621). Amino acids 662 to 853 are from the Fc domain of human IgG1 (Lewis, et

al., J. Immunol. 151:2829-2838 (1993). (†) mark the two cysteines (amino acids number 632 and 635) of the IgG hinge preceding the Fc that form the inter-chain disulfide bridges that link two Fc domains. The hexahistine tag is shown in bold/italic type (amino acids 854 to 859). (•) shows the position of the STOP codon.

FIGURE 5: The amino acid sequence of human IL-6Rα-Fc. Key: Amino acids 1 to 358 are from human IL-6Rα (Yamasaki, et al., Science 241:825-828 (1988). Note that amino acid number 2 has been changed from a Leu to a Val in order to accommodate a Kozak sequence in the coding DNA sequence. The signal peptide of IL-6Rα-Fc has been italicized (amino acids 1 to 19). The Ala-Gly bridge is shown in bold type (amino acids 359, 360). Amino acids 361 to 592 are from the Fc domain of human IgG1 (Lewis et al., J. Immunol. 151:2829-2838 (1993). (†) mark the two cysteines (amino acids number 371 and 374) of the IgG hinge preceding the Fc that form the inter-chain disulfide bridges that link two Fc domains. (•) shows the position of the STOP codon.

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FIGURE 6: The CNTF/IL-6/IL-11 receptor system. The ordered formation of the hexameric signal transducing receptor complex is depicted schematically. The cytokine associates with the Rα component to form an obligatory cytokine •Rα complex (Kd is about 5 nM). This low affinity complex next associates with the first signal transducing component, marked β1, to form a high affinity cytokine •Rα • β1 complex (Kd is about 10 pM). In the case of IL-6Rα, this component is gp130. This trimeric high affinity complex subsequently associates with another such complex. Formation of this complex results in signal transduction as it involves dimerization of two signal transducing components, marked β1 and β2 respectively (adapted from (Ward et al., J. Bio. Chem. 269:23286-23289 (1994); Stahl and Yancopoulos, J. Neurobiology 25:1454-1466 (1994); Stahl and Yancopoulos, Cell 74:587-590 (1993).

FIGURE 7: Design of heterodimeric receptor-based ligand traps for IL-6. The heterodimeric ligand trap is comprised of two interdisulfide linked proteins, gp130-Fc and IL-6Rα-Fc. The gp130-Fc•IL-6Rα-Fc complex (upper panel) is shown to mimic the high affinity cytokine•Rα•β1 complex (lower panel). The ligand trap functions as an antagonist by sequestering IL-6 and thus rendering unavailable to interact with the native receptors on IL-6-responsive cells.

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FIGURE 8. Heteromeric immunoglobulin Heavy/Light Chain Receptor Fusions. An example of a heavy/light chain receptor fusion molecule is schematically depicted. The extracellular domain of gp130 is fused to Cγ, whereas the extracellular domain of IL-6Rα is fused to the constant region of the kappa chain (κ). The inter-chain disulfide bridges are also depicted (S-S).

FIGURE 9. Amino acid sequence of gp130-Cγ1. Key: Amino acids 1 to 619 are from human gp130 (Hibi, et al., Cell 63:1149-1157 (1990). Ser-Gly bridge is shown in bold type. Amino acids 662 to 651 are from the constant region of human IgG1 (Lewis et al., J. Immunol. 151:2829-2838 (1993). (*) shows the position of the STOP codon.

FIGURE 10: Amino acid sequence of gp130 Δ 3fibro. Key: Amino acids 1 to 330 are from human gp130 (Hibi et al., Cell 63:1149-1157 (1990). Other symbols as described in Figure 9.

FIGURE 11: Amino acid sequence of J-CH1. Key: The Ser-Gly bridge is shown in bold, the J-peptide is shown in italics, the CH1 domain is underlined.

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FIGURE 12: Amino acid sequence of Cγ4. Key: The Ser-Gly bridge is shown in bold type. Amino acids 2 to 239 comprise the Cγ4 sequence.

FIGURE 13: Amino acid sequence of κ-domain. Key: The Ser-Gly bridge is shown in bold type. Amino acids 2 to 108 comprise the κ domain. The C-terminal cysteine (amino acid 108) is that involved in the disulfide bond of the κ domain with the CH1 domain of Cγ.

FIGURE 14: Amino acid sequence of λ -domain. Key: The Ser-Gly bridge is shown in bold type. Amino acids 2 to 106 comprise the λ domain (Cheung, et al., J. Virol. 66: 6714-6720 (1992). The C-terminal cysteine (amino acid 106) is that involved in the disulfide bond of the λ domain with the CH1 domain of C γ .

FIGURE 15: Amino acid sequence of the soluble IL-6Rα domain. Key:

Amino acids 1 to 358 comprise the soluble IL-6Rα domain (Yamasaki, et al., Science 241:825-828 (1988). The Ala-Gly bridge is shown in bold type.

FIGURE 16: Amino acid sequence of the soluble IL-6Rα313 domain: Key:

Amino acids 1 to 313 comprise the truncated IL-6Rα domain (IL-6Rα313).

The Thr-Gly bridge is shown in bold type.

FIGURE 17: Purification of gp130-Cγ1•IL-6Rα-κ. 4% to 12% SDS-PAGE gradient gel run under non-reducing conditions. Proteins were visualized by staining with silver. Lane 1: approximately 100 ng of material purified over Protein A Sepharose (Pharmacia). Lane 2: Molecular size standards (Amersham). Lane 3: The Protein A-purified material shown here after further purification over an IL-6 affinity chromatography step. The positions of the gp130-Cγ1 dimer [(gp130-Cγ1)2], the gp130-Cγ1 dimer

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associated with one IL-6R α - κ [(gp130-C γ 1)2•(IL-6R α - κ)1], and the gp130-C γ 1 dimer associated with two IL-6R α - κ [(gp130-C γ 1)2•(IL-6R α - κ)2] are shown, as well as the sizes for the molecular size standards in kilodaltons (200, 100, and 46).

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FIGURE 18: IL-6 dissociates slowly from the ligand trap. The dissociation rate of IL-6 from a heavy/light chain receptor-based ligand trap (gp130- $C\gamma1 \bullet IL-6R\alpha-\kappa$) was compared to that obtained with the neutralizing monoclonal antibody B-E8 (BE8 MAb).

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FIGURE 19: IL-6 can induce multimerization of the ligand trap. (A) Two different ligand traps are depicted schematically and listed according to their ability to bind protein A. gp130-Fc•IL-6R α -Fc (GF6F) binds protein A via its Fc-domains, whereas gp130-CH1•IL-6R α -K (G16K) does not bind to protein A. (B) Anti-kappa western blotting of proteins precipitated with Protein A-Sepharose from mixtures of GF6F \pm IL-6, G16K \pm IL-6, or GF6F plus G16K \pm IL-6, as marked.

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FIGURE 20: Inhibition of IL-6-dependent XG-1 cell proliferation. XG-1 cells [Zhang, et al., Blood 83:3654-3663 (1994)] were prepared for a proliferation assay by starving the cells from IL-6 for 5 hours. Assays were set up in 96-well tissue culture dishes in RPMI + 10% fetal calf serum + penicillin/streptomycin + 0.050 nM 2-mercaptoethanol + glutamine. 0.1 ml of that media was used per well. Cells were suspended at a density of 250,000 per ml at the start of the assay. 72 hours post addition of IL-6 ± ligands traps or antibodies, an MTT assay was performed as described (Panayotatos et al. Biochemistry 33:5813-5818 (1994). The different ligand traps utilized are listed.

FIGURES 21A-21D: Nucleotide sequence encoding and deduced amino acid sequence of fusion polypeptide designated 424 which is capable of binding the cytokine IL-4 to form a nonfunctional complex.

- FIGURES 22A-22D: Nucleotide sequence encoding and deduced amino acid sequence of fusion polypeptide designated 603 which is capable of binding the cytokine IL-4 to form a nonfunctional complex.
- FIGURES 23A-23D: Nucleotide sequence encoding and deduced amino acid sequence of fusion polypeptide designated 622 which is capable of binding the cytokine IL-4 to form a nonfunctional complex.

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- FIGURE 24A-24F: Nucleotide sequence encoding and deduced amino acid sequence of fusion polypeptide designated 412 which is capable of binding the cytokine IL-6 to form a nonfunctional complex.
- FIGURE 25A-25F: Nucleotide sequence encoding and deduced amino acid sequence of fusion polypeptide designated 616 which is capable of binding the cytokine IL-6 to form a nonfunctional complex.
- FIGURE 26A-26E: Nucleotide sequence encoding and deduced amino acid sequence of fusion polypeptide designated 569 which is capable of binding the cytokine IL-1 to form a nonfunctional complex.
- FIGURE 27: Shows that an IL-4 trap designated 4SC375, which is a fusion polypeptide of IL-2R γ -scb-IL4R α -Fc Δ C1, is several orders of magnitude better as an IL-4 antagonist than IL4R α Fc Δ C1 alone in the TF1 cell bioassay.
- FIGURE 28: Shows that an IL-4 trap designated 4SC375 displays, antagonistic activity in the TF1 cell bioassay equivalent to an IL-4 trap designated 4SC424 (described in Figs. 21A-21D) which is a fusion

polypeptide of IL-2R γ -IL4R α -Fc Δ C1 having the IL-2R γ component flush with the IL-4R α component.

- FIGURE 29: Shows that the IL6 trap (6SC412 IL6R-scb-gpx-FcΔC1) described in Figs. 24A-24F is a better antagonist of IL-6 in the XG1 bioassay than the neutralizing monoclonal antibody to human IL-6 BE8.
- FIGURE 30: Shows that the trap 1SC569 (described in Figs. 26A-26E) is able to antagonize the effects of IL-1 and block the IL-6 production from MRC 5 cells upon treatment with IL-1.
 - FIGURE 31A-31G: The nucleotide and encoded amino acid sequence of the IL-4Rα.IL-13Rα1.Fc single chain trap construct is set forth.
- FIGURE 32A-32G: The nucleotide and encoded amino acid sequence of the IL-13Rα1.IL-4Rα.Fc single chain trap construct is set forth.
- FIGURE 33: Blocking of IL-13 by IL-4Rα.IL-13Rα1.Fc and IL-13Rα1.IL-4Rα.Fc. Addition of either IL-4Rα.IL-13Rα1.Fc or IL-13Rα1.IL-4Rα.Fc trap at a concentration of 10nM blocks IL-13-induced growth up to ~2nM. At an IL-13 concentration of ~4-5 nM the growth of TF1 cells is inhibited by 50%.
- FIGURE 34: Blocking of IL-4 by IL-4Rα.IL-13Rα1.Fc and IL-13Rα1.IL-4Rα.Fc.

 Addition of either IL-4Rα.IL-13Rα1.Fc or IL-13Rα1.IL-4Rα.Fc at a

 concentration of 10nM blocks IL-4-induced growth up to ~1nM. At an IL-4 concentration of ~3-4 nM the growth of TF1 cells is inhibited by 50%.
- FIGURE 35: Human IL-1 trap blocks the in vivo effects of exogenously administered huIL-1. BALB/c mice were given subcutaneous injection of huIL-1 (0.3 µg/kg) at time 0. Twenty-four hours prior to huIL-1 injection, the animals were pre-treated with either vehicle or 150-fold molar excess

of huIL-1 trap. Two hours prior to sacrifice (26 hrs), the mice were rechallenged with a second injection of huIL-1 (0.3 μ g/kg, s.c.). Blood samples were collected at various time points and sera were assayed for IL-1 levels (expressed as mean +/- SEM; n=5 per group).

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FIGURE 36A & FIGURE 36B: Human IL-4 trap antagonizes the effects of human IL-4 in monkeys. Figure 36A: Cynomologus monkeys were treated in three parts as indicated. Human IL-4 (25 µg/kg) was injected subcutaneously twice daily for 4 days and human IL-4 trap (8 mg/ml) and vehicle were given intravenously daily for 5 days, beginning 1 day prior to human IL-4 administration. Plasma was collected daily and assayed for MCP-1 levels. Results were expressed as mean +/- SEM; n=4. (ANOVA p<0.0007; Tukey-Kramer: Part 2 vs. Part 1, p,0.05; Part 2 vs. Part 3, p,0.05; Part 1 vs. Part 3, not significant.) Figure 36B: Cynomologus monkeys were treated in three parts as indicated. Human IL-4 (25 µg/kg) was injected subcutaneously twice daily for 4 days and human IL-4 trap (8 mg/ml) and vehicle were given intravenously daily for 5 days, beginning 1 day prior to human IL-4 administration. Whole blood was collected daily for flow cytometry analysis for CD16. Results were expressed as mean +/- SEM; n=4. (ANOVA p<0.042; Tukey-Kramer: Part 2 vs. Part 1, p<0.05; Part 2 vs. Part 3 and Part 1 vs. Part 3, not significant.)

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FIGURE 37: Murine IL-4 trap partially prevented IL-4-mediated IgE increase in mice. BALB/C mice injected with anti-mouse IgD

25 (100μl/mouse, s.c.) were randomly divided into 3 groups, each received (on days 3-5) either vehicle, murine IL-4 trap (1 mg/kg, s.c.), or a monoclonal antibody to mouse IL-4 (1 mg/kg, s.c.). Sera were collected at various time points and assayed for IgE levels. Results were expressed as mean+/-SEM (n=5 per group). (ANOVA p=0.0002; Tukey-Kramer: vehicle vs. IL-4 trap, p<0.01; vehicle vs. IL-4 antibody, p<0.001; IL-4 trap vs. IL-4 antibody, not significant).

DETAILED DESCRIPTION OF THE INVENTION

The present invention provides an isolated nucleic acid molecule encoding a fusion polypeptide capable of binding a cytokine to form a nonfunctional complex comprising:

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- a) a nucleotide sequence encoding a first fusion polypeptide component comprising the amino acid sequence of the cytokine binding portion of the extracellular domain of the specificity determining component of the cytokine's receptor;
- b) a nucleotide sequence encoding a second fusion polypeptide component comprising the amino acid sequence of the cytokine binding portion of the extracellular domain of the signal transducing component of the cytokine's receptor; and
 - c) a nucleotide sequence encoding a third fusion polypeptide component comprising the amino acid sequence of a multimerizing component.

By "cytokine binding portion" what is meant is the minimal portion of the extracellular domain necessary to bind the cytokine. It is accepted by those of skill in the art that a defining characteristic of a cytokine receptor is the presence of the two fibronectin-like domains that contain canonical cysteines and of the WSXWS box (Bazan, J.F., 1990, PNAS 87: 6934-6938). Sequences encoding the extracellular domains of the binding component of the cytokine's receptor and of the signal transducing component of the cytokine's receptor may also be used to create the fusion polypeptide of the invention. Similarly, longer sequences encoding larger portions of the components of the cytokine's receptor may be used. However, it is contemplated that fragments smaller than the extracellular domain will function to bind the cytokine and therefore, the invention contemplates fusion polypeptides comprising the minimal portion of the extracellular domain necessary to bind the cytokine as the cytokine binding portion.

The invention comprises a "specificity determining component" of a cytokine's receptor and a "signal transducing component" of the cytokine's receptor. Regardless of the nomenclature used to designate a particular component or subunit of a cytokine receptor, one skilled in the art would recognize which component or subunit of a receptor is responsible for determining the cellular target of the cytokine, and thus would know which component constitutes the "specificity determining component."

Similarly, regardless of the nomenclature used, one of skill in the art would know which component or subunit of a receptor would constitute the "signal transducing component." As used herein, the "signal transducing component" is a component of the native receptor which is not the specificity determining component and which does not bind or weakly binds the cytokine in the absence of the specificity determining component. In the native receptor, the "signal transducing component" may participate in signaling.

For example, while some cytokine receptors have components designated α and β , the IL-4 receptor has a signal transducing component referred to as IL-2R γ . However, regardless of what name is associated with that component, one skilled in the art would know which component of the IL-4 receptor is the signal transducing component. Thus to practice the present invention and create a high affinity trap for IL-4, one of skill in the art would create an isolated nucleic acid comprising a nucleotide sequence encoding a first fusion polypeptide component comprising the amino acid sequence of the cytokine binding portion of the extracellular domain of the specificity determining component of the IL-4 receptor (IL-4R α); a nucleotide sequence encoding a second fusion polypeptide component comprising the amino acid sequence of the cytokine binding portion of the extracellular domain of the signal transducing component of the IL-4 receptor (IL-2R γ); and a nucleotide sequence encoding a third fusion polypeptide component comprising the amino acid sequence of a

multimerizing component (for example, an Fc domain of IgG) to create a high affinity trap for IL-4.

Some further examples of the receptor components that may be used to

5 prepare cytokine antagonists according to the invention are set forth in
Table 1. The Table 1 sets forth, by way of example but not by way of
limitation, some of the varied nomenclature used in the scientific
literature to describe those components which function as specificity
determining components and those which function as signal transducing

10 components of certain cytokine receptors.

TABLE

Cytokine	Specificity determining Component	Signal transducing Component
Interleukin-1 (IL-1)	Type I IL-1R (ref. 8) Type II IL-1R (ref. 8) IL-1RI (ref. 11) IL-1RII (ref. 11)	IL-1R AcP (refs. 8, 11).
Interleukin-2 (IL-2)	α -subunit (ref. 2) α -chain (ref. 3) IL-2R α (ref. 1)	β-chain (ref. 3) β-subunit (ref. 2) γ-chain (ref. 3) IL-2Rβ (refs. 1, 10)
Interleukin-3 (IL-3)	IL-3R α (ref. 1) α -subunit (ref. 2) α -receptor component (ref. 5)	IL-2Ry (refs. 1, 10) $\beta_{c} \text{ (ref. 1)}$ $\beta\text{-subunit (ref. 2)}$ $\beta\text{-chain (ref. 3)}$ $\beta\text{-recentor component (ref. 5)}$
Interleukin-4 (IL-4)	IL-4R (ref. 1)	
Interleukin-5 (IL-5)	IL-5R α (ref. 1) α -subunit (ref. 2) α -receptor component (ref. 5)	β _c (ref. 1) β-subunit (ref. 2) β-chain (ref. 3) β-receptor component (ref. 5)

TABLE 1 (CONT'D)

nponent Signal transducing Component	β-receptor component (ref. 5) β-subunit (ref. 2) β-chain (ref. 3) β _c (ref. 1) GMRβ (refs. 1, 2)	gp130 (refs. 1, 3) β- receptor component (ref. 5)	gp130 (ref. 4)	IL-2R β (ref. 10) IL-2R γ (ref. 10)	AF-1 (ref. 7) IFN-γR2 (ref. 7)	Type I (refs. 6, 9)
Specificity determining Compone	α -receptor component (ref. 5) α -subunit (ref. 2) GMR α (refs. 1, 2)	LIFBP (ref. 1) α-receptor component (ref. 5)	α-chain (ref. 4) NR1 (ref. 4)	IL-15 $R\alpha$ (ref. 10)	IFN-γR (ref. 7) IFN-γR1 (ref. 7)	Type II (refs. 6, 9)
Cytokine	Granulocyte macrophage- colony stimulating factor (GM-CSF)	Leukemia inhibitory factor (LIF)	Interleukin-11 (IL-11)	Interleukin-15 (IL-15)	Interferon-γ (IFNγ)	TGFB

Only a few of the multitude of references are cited in Table 1, and they are set forth as follows:

- 1. Sato and Miyajima, Current Opinions in Cell Biology 6: 174-179
- 5 (1994) See page 176, lines 9-16;
 - 2. Miyajima, et al., Annual Review of Immunology 10: 295-331 (1992) See page 295, line 4 to page 296, line 1; page 305, last paragraph;
 - 3. Kondo, et al, Science 262: 1874-1877 (1993) See page 1874, cols. 1 & 2;
 - 4. Hilton, et al, EMBO Journal 13: 4765-4775 (1994) See page 4766, col.
- 10 1, lines 20 24;
 - 5. Stahl and Yancopoulos, Cell 74: 587-590 (1993) See page 587, column 2, lines 15-22;
 - 6. Bassing, et al, Journal of Biological Chemistry 269: 14861-14864 (1994)
 See page 14861, col. 2, lines 1-9 and 21-28;
- 7. Kotenko, et al, Journal of Biological Science 270: 20915-20921 (1995) See page 20915, lines 1-5 of the abstract;
 - 8. Greenfeder, et al., Journal of Biological Chemistry 270: 13757-13765 (1995) See page 13757, col. 1, line 6 to col. 2, line 3 and col. 2, lines 10-12; page 13764, col. 2, last 3 lines and page 13765, col. 1, lines 1-7;
- 20 9. Lebrun and Vale, Molecular Cell Biology 17: 1682-1691 (1997) See page 1682, Abstract lines 2-6;
 - 10. Kennedy and Park, Journal of Clinical Immunology 16: 134-143 (1996) See page 134, lines 1-7 of the abstract; page 136, col 2., lines 1-5;
 - 11. Wesche, et al., Journal of Biological Chemistry 272: 7727-7731 (1997)
- 25 See page 7731, lines 20-26.

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Kotenko, et al. recently identified the IL-10R2 (IL-10Rβ) chain which is reported to serve as an accessory chain that is essential for the active IL-10 receptor complex and for initiating IL-10 induced signal transduction events (S.V. Kotenko, et al., The EMBO Journal, 1997, Vol. 16: 5894-5903). Additional cytokines and their receptors are described in Appendix II, page A:9 of Immunobiology, The Immune System In Health and Disease, 2nd

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In preparing the nucleic acid sequence encoding the fusion polypeptide of the invention, the first, second, and third components of the fusion polypeptide are encoded in a single strand of nucleotides which, when expressed by a host vector system, produces a monomeric species of the fusion polypeptide. The monomers thus expressed then multimerize due to the interactions between the multimerizing components (the third fusion polypeptide components). Producing the fusion polypeptides in this manner avoids the need for purification of heterodimeric mixtures that would result if the first and second components were produced as separate molecules and then multimerized. For example, U.S. Patent No. 5,470,952 issued November 28, 1995 describes the production of heterodimeric proteins that function as CNTF or IL-6 antagonists. The 15 heterodimers are purified from cell lines cotransfected with the appropriate alpha (α) and beta (β) components. Heterodimers are then separated from homodimers using methods such as passive elution from preparative, nondenaturing polyacrylamide gels or by using high pressure cation exchange chromatography. The need for this purification step is 20 avoided by the methods of the present invention.

In addition, PCT International Application WO 96/11213 published 18 April 1996 entitled Dimeric IL-4 Inhibitors states that the applicant has prepared homodimers in which two IL-4 receptors are bound by a polymeric spacer and has prepared heterodimers in which an IL-4 receptor is linked by a polymeric spacer to an IL-2 receptor gamma chain. The polymeric spacer described is polyethylene glycol (PEG). The two receptor components, IL-4R and IL-2Rgamma are separately expressed and purified. Pegylated homodimers and heterodimers are then produced by joining the components together using bi-functional PEG reagents. It is an advantage

of the present invention that it avoids the need for such time consuming and costly purification and pegylation steps.

In one embodiment of the invention, the nucleotide sequence encoding the first component is upstream of the nucleotide sequence encoding the second component. In another embodiment of the invention, the nucleotide sequence encoding the first component is downstream of the nucleotide sequence encoding the second component. Further embodiments of the invention may be prepared in which the order of the first, second and third fusion polypeptide components are rearranged. For example, if the nucleotide sequence encoding the first component is designated 1, the nucleotide sequence encoding the second component is designated 2, and the nucleotide sequence of the third component is designated 3, then the order of the components in the isolated nucleic acid of the invention as read from 5' to 3' may be any of the following six combinations: 1,2,3; 1,3,2; 2,1,3; 2,3,1; 3,1,2; or 3,2,1.

In further embodiments of the invention, the cytokine bound by the fusion polypeptide may be a member of the hematopoietin family of cytokines selected from the group consisting of interleukin-2, interleukin-3, interleukin-4, interleukin-5, interleukin-6, interleukin-7, interleukin-9, interleukin-11, interleukin-13, interleukin-15, granulocyte macrophage colony stimulating factor, oncostatin M, leukemia inhibitory factor, and cardiotrophin-1.

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In additional embodiments of the invention, the cytokine bound by the fusion polypeptide may be a member of the interferon family of cytokines selected from the group consisting of IFN-gamma, IFN-alpha, and IFN-beta.

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In additional embodiments of the invention, the cytokine bound by the fusion polypeptide may be a member of the immunoglobulin superfamily

of cytokines selected from the group consisting of B7.1 (CD80) and B7.2 (B70).

In still further embodiments of the invention, the cytokine bound by the fusion polypeptide may be a member of the TNF family of cytokines selected from the group consisting of TNF-alpha, TNF-beta, LT-beta, CD40 ligand, Fas ligand, CD 27 ligand, CD 30 ligand, and 4-1BBL.

In additional embodiments of the invention, the cytokine bound by the fusion polypeptide may be a cytokine selected from the group consisting of interleukin-1, interleukin-10, interleukin-12, interleukin-14, interleukin-18, and MIF.

Because specificity determination and signal transduction occurs by a

similar mechanism in the TGF-β/BMP family of cytokines (See D.

Kingsley, Genes & Development, 1994, 8: 133-146; J. Wrana, Miner

Electrolyte Metab, 24: 120-130 (1998); R. Derynck and X. Feng, Biochimica et

Biophysica Acta 1333 (1997) F105-F150; and J. Massague and F. Weis-Garcia,

"Serine/threonine Kinase Receptors: Mediators of Transforming Growth

Factor Beta Family Signals" In Cancer Surveys, Vol. 27: Cell Signaling,

1996, Imperial Cancer Research Fund) the present invention may be used

to produce high affinity antagonists for cytokines that are members of the

TGF-β/BMP family.

Therefore, in additional embodiments of the invention, the cytokine bound by the fusion polypeptide may be a member of the TGF-β/BMP family selected from the group consisting of TGF-β1, TGF-β2, TGF-β3, BMP-2, BMP-3a, BMP-3b, BMP-4, BMP-5, BMP-6, BMP-7, BMP-8a, BMP-8b, BMP-9, BMP-10, BMP-11, BMP-15, BMP-16, endometrial bleeding associated factor (EBAF), growth differentiation factor-1 (GDF-1), GDF-2, GDF-3, GDF-5, GDF-6, GDF-7, GDF-8, GDF-9, GDF-12, GDF-14, mullerian

inhibiting substance (MIS), activin-1, activin-2, activin-3, activin-4, and activin-5.

In alternative embodiments of the invention, the specificity determining component, the signal transducing component, or both, may be substituted 5 for by a single chain Fv. A single chain Fv (scFv) is a truncated Fab having only the V region of a heavy chain linked by a stretch of synthetic peptide to a V region of a light chain. See, for example, US Patent Nos. 5,565,332; 5,733,743; 5,837,242; 5,858,657; and 5,871,907 assigned to Cambridge Antibody Technology Limited incorporated by reference herein. Thus the 10 present invention contemplates, for example, an isolated nucleic acid molecule encoding a fusion polypeptide capable of binding a cytokine to form a nonfunctional complex comprising a nucleotide sequence encoding a first fusion polypeptide component comprising the amino acid sequence of the cytokine binding portion of the extracellular domain of the 15 specificity determining component of the cytokine's receptor; a nucleotide sequence encoding a second fusion polypeptide component comprising the amino acid sequence of an scFv capable of binding the cytokine at a site different from the site at which the cytokine binding portion of the extracellular domain of the specificity determining component of the cytokine's receptor binds; and a nucleotide sequence encoding a third fusion polypeptide component comprising the amino acid sequence of a multimerizing component. Alternatively, the specificity determining component may be substituted for by a scFv that binds to a site on the cytokine different from the site at which the signal transducing 25 component binds. Thus the invention contemplates an isolated nucleic acid molecule encoding a fusion polypeptide capable of binding a cytokine to form a nonfunctional complex comprising a nucleotide sequence encoding a first fusion polypeptide component comprising the amino acid sequence of a scFv that binds to a site on the cytokine different from the 30 site at which the cytokine binding portion of the extracellular domain of the signal transducing component of the cytokine's receptor binds; a nucleotide sequence encoding a second fusion polypeptide component

comprising the amino acid sequence of the cytokine binding portion of the extracellular domain of the signal transducing component of the cytokine's receptor; and a nucleotide sequence encoding a third fusion polypeptide component comprising the amino acid sequence of a multimerizing component.

In another embodiment, the invention contemplates an isolated nucleic acid molecule encoding a fusion polypeptide capable of binding a cytokine to form a nonfunctional complex comprising a nucleotide sequence encoding a first fusion polypeptide component comprising the amino acid sequence of a first scFv that binds to a site on the cytokine; a nucleotide sequence encoding a second fusion polypeptide component comprising the amino acid sequence a second scFv that binds to a site on the cytokine different from the site at which the first scFv binds; and a nucleotide sequence encoding a third fusion polypeptide component comprising the amino acid sequence of a multimerizing component.

In all of the above described embodiments comprising scFv's, the invention also contemplates embodiments in which the nucleotide sequence encoding the first component is upstream of the nucleotide sequence encoding the second component; embodiments in which the nucleotide sequence encoding the first component is downstream of the nucleotide sequence encoding the second component; and further embodiments of the invention in which the order of the first, second and third fusion polypeptide components is rearranged. For example, if the nucleotide sequence encoding the first component is designated 1, the nucleotide sequence encoding the second component is designated 2, and the nucleotide sequence of the third component is designated 3, then the order of the components in the isolated nucleic acid of the invention as read from 5' to 3' may be any of the following six combinations: 1,2,3; 1,3,2; 2,1,3; 2,3,1; 3,1,2; or 3,2,1.

In preferred embodiments of the invention, the multimerizing component comprises an immunoglobulin derived domain. More specifically, the immunoglobulin derived domain may be selected from the group consisting of the Fc domain of IgG, the heavy chain of IgG, and the light chain of IgG. In another embodiment, the multimerizing component may be an Fc domain from which the first five amino acids (including a cysteine) have been removed to produce a multimerizing component referred to as $Fc(\Delta C1)$. Alternatively, the multimerizing component may be an Fc domain in which a cysteine within the first five amino acids has been substituted for by another amino acid such as, for example, serine or alanine.

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The present invention also provides for fusion polypeptides encoded by the isolated nucleic acid molecules of the invention. Preferably, the fusion polypeptides are in multimeric form, due to the function of the third multimerizing component. In a preferred embodiment, the multimer is a dimer. Suitable multimerizing components are sequences encoding an immunoglobulin heavy chain hinge region (Takahashi et al., 1982, Cell 29:671-679); immunoglobulin gene sequences, and portions thereof. In a preferred embodiment of the invention, immunoglobulin gene sequences, especially one encoding the Fc domain, are used to encode the third multimerizing component.

The present invention also contemplates a vector which comprises the nucleic acid molecule of the invention as described herein.

Also provided is an expression vector comprising a nucleic acid molecule of the invention as described herein, wherein the nucleic acid molecule is operatively linked to an expression control sequence. Also provided is a host-vector system for the production of a fusion polypeptide which comprises the expression vector of the invention which has been introduced into a host cell suitable for expression of the fusion

polypeptide. The suitable host cell may be a bacterial cell such as <u>E. coli</u>, a yeast cell, such as <u>Pichia pastoris</u>, an insect cell, such as <u>Spodoptera</u> frugiperda, or a mammalian cell, such as a COS, CHO, 293, BHK or NS0 cell.

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The present invention also provides for methods of producing the fusion polypeptides of the invention by growing cells of the host-vector systems described herein, under conditions permitting production of the fusion polypeptide and recovering the fusion polypeptide so produced.

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The present invention provides novel antagonists which are based on receptor components that are shared by cytokines such as the CNTF family of cytokines.

The invention described herein contemplates the production of antagonists to any cytokine that utilizes an α specificity determining component which, when combined with the cytokine, binds to a first β signal transducing component to form a nonfunctional intermediate which then binds to a second β signal transducing component causing β-receptor dimerization and consequent signal transduction. According to the invention, the soluble α specificity determining component of the receptor (sRα) and the extracellular domain of the first β signal transducing component of the cytokine receptor (β1) are combined to form heterodimers (sRα:β1) that act as antagonists to the cytokine by binding the cytokine to form a nonfunctional complex.

As described in Example 1, CNTF and IL-6 share the $\beta 1$ receptor component gp130. The fact that CNTF forms an intermediate with CNTFR α and gp130 can be demonstrated (Example 1) in cells lacking LIFR β , where the complex of CNTF and CNTFR α binds gp130, and

prevents homodimerization of gp130 by IL-6 and IL-6Rα, thereby blocking signal transduction. These studies provide the basis for the development of the IL-6 antagonists described herein, as they show that if, in the presence of a ligand, a nonfunctional intermediate complex, consisting of the ligand, its α receptor component and its β1 receptor component, can be formed, it will effectively block the action of the ligand. Other cytokines may use other β1 receptor components, such as LIFRβ, which may also be used to produce antagonists according to the present invention.

- Thus for example, in one embodiment of the invention, effective antagonists of IL-6 or CNTF consist of heterodimers of the extracellular domains of the α specificity determining components of their receptors (sIL-6Rα and sCNTFRα, respectively) and the extracellular domain of gp130. The resultant heterodimers, which are referred to hereinafter as sIL-6Rα:β1 and sCNTFRα:β1, respectively, function as high-affinity traps for IL-6 or CNTF, respectively, thus rendering the cytokine inaccessible to form a signal transducing complex with the native membrane-bound forms of their receptors.
- Although soluble ligand binding domains from the extracellular portion of receptors have proven to be somewhat effective as traps for their ligands and thus act as antagonists [Bargetzi, et al., Cancer Res. 53:4010-4013 (1993); et al., Proc. Natl. Acad. Sci. USA 89: 8616-8620 (1992); Mohler, et al., J. Immunol. 151: 1548-1561 (1993); Narazaki, et al., Blood 82: 1120-1126 (1993)], the IL-6 and CNTF receptors are unusual in that the α receptor components constitute ligand binding domains that, in concert with their ligands, function effectively in soluble form as receptor agonists [Davis, et al. Science 259:1736-1739 (1993); Taga, et al., Cell 58: 573-581 (1989)]. The sRα:β1 heterodimers prepared according to the present invention provide effective traps for their ligands, binding these ligands with affinities in the picomolar range (based on binding studies for CNTF to PC12D cells)

without creating functional intermediates. The technology described herein may be applied to develop a cytokine trap for any cytokine that utilizes an α -component that confers specificity, as well as a β component which, when bound to the α -specificity component, has a higher affinity for the cytokine than either component alone. Accordingly, antagonists according to the invention include antagonists of interleukins 1 through 5 [IL-1, Greenfeder, et al. J. Biol. Chem. 270:13757-13765 (1995); Guo, et al. J. Biol. Chem. 270:27562-27568 (1995)], IL-2; [Taniguchi, et al. European Patent Nos. 0386289-A and 0386304-A (1990); Takeshita, et al. Science 257:379-382 (1992)]; IL-3; [Kitamura, et al. Cell 66:1165-1174 (1991)], IL-4; [Idzerda, et al. J. Exp. Med. 171:861-873 (1990)], IL-5; [Taverneir, et al. Cell 66:1175-1184 (1991)], IL-11 [(Cherel, et al. Direct Submission to EMBL/GenBank/DDB] databases; accession No. Z38102)], interleukin 15 [IL-15; Hemar, et al. J. Cell Biol. 1295:55-64 (1995); Taniguchi, et al. European Patent Nos. 0386289-A and 0386304-A (1990); Takeshita, et al. Science 257:379-382 (1992)], granulocyte-macrophage colony stimulating factor [GM-CSF; Hayashida, et al. Proc. Natl. Acad. Sci. U.S.A. 97:9655-9659 (1990)], LIF, gamma interferon [IFNy; Aguet, et al. Cell 55:273-280 (1988); Soh, et al. Cell 76:793-802 (1994)], and transforming growth factor beta [TGFβ; Inagaki, et al. Proc. Natl. Acad. Sci. USA 90:5359-5363 (1993)].

The α and β receptor extracellular domains may be prepared using methods known to those skilled in the art. The CNTFR α receptor has been cloned, sequenced and expressed [Davis, et al. (1991) Science 253:59-63 which is incorporated by reference in its entirety herein]. The cloning of LIFR β and gp130 are described in Gearing et al. in EMBO J. 10:2839-2848 (1991), Hibi, et al. Cell 63:1149-1157 (1990) and in published PCT application WO 93/10151 published May 27, 1993, all of which are incorporated by reference in their entirety herein.

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The receptor molecules useful for practicing the present invention may be prepared by cloning and expression in a prokaryotic or eukaryotic expression system. The recombinant receptor gene may be expressed and purified utilizing any number of methods. The gene encoding the factor may be subcloned into a bacterial expression vector, such as for example, but not by way of limitation, pCP110.

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The recombinant factors may be purified by any technique which allows for the subsequent formation of a stable, biologically active protein. For example, and not by way of limitation, the factors may be recovered from cells either as soluble proteins or as inclusion bodies, from which they may be extracted quantitatively by 8M guanidinium hydrochloride and dialysis. In order to further purify the factors, conventional ion exchange chromatography, hydrophobic interaction chromatography, reverse phase chromatography or gel filtration may be used.

The sR α : β heterodimeric receptors may be engineered using known fusion regions, as described in published PCT application WO 93/10151 published May 27, 1993 entitled "Receptor for Oncostatin M and Leukemia Inhibitory Factor" which describes production of β receptor heterodimers, or they may be prepared by crosslinking of extracellular domains by chemical means. The domains utilized may consist of the entire extracellular domain of the α and β components, or they may consist of mutants or fragments thereof that maintain the ability to form a complex with its ligand and other components in the sR α : β 1 complex. For example, as described below in Example 4, IL-6 antagonists have been prepared using gp130 that is lacking its three fibronectin-like domains.

In one embodiment of the invention, the extracellular domains are engineered using leucine zippers. The leucine zipper domains of the human transcription factors c-jun and c-fos have been shown to form stable heterodimers [Busch and Sassone-Corsi, Trends Genetics 6: 36-40]

(1990); Gentz, et al., Science 243: 1695-1699 (1989)] with a 1:1 stoichiometry. Although jun-jun homodimers have also been shown to form, they are about 1000-fold less stable than jun-fos heterodimers. Fos-fos homodimers have not been detected.

The leucine zipper domain of either c-jun or c-fos are fused in frame at the C-terminus of the soluble or extracellular domains of the above mentioned receptor components by genetically engineering chimeric genes. The fusions may be direct or they may employ a flexible linker domain, such as the hinge region of human IgG, or polypeptide linkers consisting of small amino acids such as glycine, serine, threonine or alanine, at various lengths and combinations. Additionally, the chimeric proteins may be tagged by His-His-His-His-His-His (His6),[SEQ. ID NO. 1] to allow rapid purification by metal-chelate chromatography, and/or by epitopes to which antibodies are available, to allow for detection on western blots, immunoprecipitation, or activity depletion/blocking in bioassays.

In another embodiment, as described below in Example 3, the sRa:\(\beta\)1 heterodimer is prepared using a similar method, but using the Fc-domain of human IgG1 [Aruffo, et al., Cell 67:35-44 (1991)]. In contrast to the latter, formation of heterodimers must be biochemically achieved, as chimeric molecules carrying the Fc-domain will be expressed as disulfide-linked homodimers. Thus, homodimers may be reduced under conditions that favor the disruption of inter-chain disulfides but do not effect intra-chain disulfides. Then monomers with different extracellular portions are mixed in equimolar amounts and oxidized to form a mixture of homoand heterodimers. The components of this mixture are separated by chromatographic techniques. Alternatively, the formation of this type of heterodimers may be biased by genetically engineering and expressing molecules that consist of the soluble or extracellular portion of the receptor components followed by the Fc-domain of hIgG, followed by

either the c-jun or the c-fos leucine zippers described above [Kostelny, et al., J. Immunol. 148: 1547-1553 (1992)]. Since these leucine zippers form predominately heterodimers, they may be used to drive formation of the heterodimers where desired. As for the chimeric proteins described using leucine zippers, these may also be tagged with metal chelates or an epitope. This tagged domain can be used for rapid purification by metal-chelate chromatography, and/or by antibodies, to allow for detection on western blots, immunoprecipitation, or activity depletion/blocking in bioassays.

In additional embodiments, heterodimers may be prepared using other 10 immunoglobulin derived domains that drive the formation of dimers. Such domains include, for example, the heavy chains of IgG (Cy1 and Cy4), as well as the constant regions of kappa (κ) and lambda (λ) light chains of human immunoglobulins. The heterodimerization of Cy with the light 15 chain occurs between the CH1 domain of Cy and the constant region of the light chain (CL), and is stabilized by covalent linking of the two domains via a single disulfide bridge. Accordingly, as described in Example 4, constructs may be prepared using these immunoglobulin domains. Alternatively, the immunoglobulin domains include domains that may be derived from T cell receptor components which drive dimerization. 20 In another embodiment of the invention, the sRα:β1 heterodimers are prepared by expression as chimeric molecules utilizing flexible linker loops. A DNA construct encoding the chimeric protein is designed such that it expresses two soluble or extracellular domains fused together in 25 tandem ("head to head") by a flexible loop. This loop may be entirely artificial (e.g. polyglycine repeats interrupted by serine or threonine at a certain interval) or "borrowed" from naturally occurring proteins (e.g. the hinge region of hIgG). Molecules may be engineered in which the order of the soluble or extracellular domains fused is switched (e.g.

30 sIL6Rα/loop/sgp130 or sgp130/loop/sIL-6Rα) and/or in which the length

and composition of the loop is varied, to allow for selection of molecules with desired characteristics.

Alternatively, the heterodimers made according to the present invention may be purified from cell lines cotransfected with the appropriate α and β components. Heterodimers may be separated from homodimers using methods available to those skilled in the art. For example, limited quantities of heterodimers may be recovered by passive elution from preparative, nondenaturing polyacrylamide gels. Alternatively, heterodimers may be purified using high pressure cation exchange chromatography. Excellent purification has been obtained using a Mono S cation exchange column.

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In addition to sRα:β1 heterodimers that act as antagonists by binding free CNTF or IL-6, the present invention also contemplates the use of engineered, mutated versions of IL-6 with novel properties that allow it to bind to IL-6Ra and a single gp130 molecule, but fail to engage the second gp130 to complete β component homodimerization, and thus act as an effective IL-6 antagonist on any IL-6 responsive cell. Our model for the structure of the IL-6 and CNTF receptor complexes indicates that these cytokines have distinct sites for binding the α , β 1, and β 2 receptor components [Stahl and Yancopoulos, Cell 74: 587-590 (1993)]. Mutations of critical amino acid residues comprising each of these sites gives rise to novel molecules which have the desired antagonistic properties. Ablation of the $\beta 1$ site would give a molecule which could still bind to the α receptor component but not the $\beta1$ component, and thereby comprise an antagonist with nanomolar affinity. Mutations of critical amino acid residues comprising the β 2 site of IL-6 (IL-6 β 2-) would give a molecule that would bind to IL-6Rα and the first gp130 monomer, but fail to engage the second gp130 and thus be functionally inactive. Similarly, mutations of

the CNTF $\beta 2$ site would give a molecule (CNTF $\beta 2$ -) that would bind CNTFR α and gp130, but fail to engage LIFR β , thereby antagonizing CNTF action by forming the non-functional $\beta 1$ intermediate. Based on the binding results described above where CNTF forms the $\beta 1$ intermediate with high affinity, both CNTF $\beta 2$ - and IL-6 $\beta 2$ - would constitute antagonists with affinity in the range of 10 pM.

A variety of means are used to generate and identify mutations of IL-6 or CNTF that have the desired properties. Random mutagenesis by standard methods of the DNA encoding IL-6 or CNTF may be used, followed by analysis of the collection of products to identify mutated cytokines having the desired novel properties as outlined below. Mutagenesis by genetic engineering has been used extensively in order to elucidate the structural organization of functional domains of recombinant proteins. Several different approaches have been described in the literature for carrying out deletion or substitution mutagenesis. The most successful appear to be alanine scanning mutagenesis [Cunningham and Wells (1989), Science 244: 1081-1085] and homolog-scanning mutagenesis [Cunningham, et al., (1989), Science 243:1330-1336].

Targeted mutagenesis of the IL-6 or CNTF nucleic acid sequences using such methods can be used to generate CNTF β 2- or IL-6 β 2- candidates. The choice of regions appropriate for targeted mutagenesis is done systematically, or determined from studies whereby panels of monoclonal antibodies against each factor are used to map regions of the cytokine that might be exposed after binding of the cytokine to the α receptor component alone, or to the $\alpha\beta1$ heterodimeric soluble receptors described above. Similarly, chemical modification or limited proteolysis of the cytokine alone or in a complex bound to the α receptor component or the $\alpha\beta1$ heterodimeric soluble receptors described above, followed by analysis

of the protected and exposed regions could reveal potential $\beta 2$ binding sites.

Assays for identifying CNTF or IL-6 mutants with the desired properties involve the ability to block with high affinity the action of IL-6 or CNTF on appropriately responsive cell lines [Davis, et al., Science 259: 1736-1739 (1993); Murakami, et al., Proc. Natl. Acad. Sci. USA 88: 11349-11353 (1991)]. Such assays include cell proliferation, survival, or DNA synthesis driven by CNTF or IL-6, or the construction of cell lines where binding of factor induces production of reporters such as CAT or β-galactosidase [Savino, et al., Proc. Natl. Acad. Sci. USA 90: 4067-4071 (1993)].

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Alternatively, the properties of various mutants may be assessed with a receptor-based assay. One such assay consists of screening mutants for their ability to bind the sRα:β1 receptor heterodimers described above using epitope-tagged [Davis et al., Science 253: 59-63 (1991)] sRα:β1 reagents. Furthermore, one can probe for the presence or absence of the β2 site by assessing whether an epitope-tagged soluble β2 reagent will bind to the cytokine in the presence of the β1 heterodimer. For example, CNTF only binds to LIFRβ (the β2 component) in the presence of both CNTFRα and gp130 [Davis, et al. Science 260: 1805-1808 (1993); Stahl, et al. J. Biol. Chem. 268: 7628-7631 (1993)]. Thus a soluble LIFRβ reagent would only bind to CNTF in the presence of the soluble sRα:β1 dimer sCNTFRα:β1. For IL-6, the sRα:β1 reagent would be IL-6Rα:β1, and the probe for the β2 site would be epitope-tagged sgp130. Thus β2- mutants of CNTF would be identified as those that bound the sRα:β1 reagent, demonstrating that the α and β1 site of the cytokine were intact, yet failed to bind the β2 reagent.

In addition, the present invention provides for methods of detecting or measuring the activity of potential β 2- mutants by measuring the phosphorylation of a β -receptor component or a signal transduction component selected from the group consisting of Jak1, Jak2 and Tyk2 or any other signal transduction component, such as the CLIPs, that are determined to be phosphorylated in response to a member of the CNTF family of cytokines.

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A cell that expresses the signal transduction component(s) described herein may either do so naturally or be genetically engineered to do so. For example, Jak1 and Tyk-2-encoding nucleic acid sequences obtained as described in Velazquez, et al., Cell, Vol. 70:313-322 (1992), may be introduced into a cell by transduction, transfection, microinjection, electroporation, via a transgenic animal, etc., using any known method known in the art.

According to the invention, cells are exposed to a potential antagonist and the tyrosine phosphorylation of either the β -component(s) or the signal transduction component(s) are compared to the tyrosine phosphorylation of the same component(s) in the absence of the potential antagonist. In another embodiment of the invention, the tyrosine phosphorylation that results from contacting the above cells with the potential antagonist is compared to the tyrosine phosphorylation of the same cells exposed to the parental CNTF family member. In such assays, the cell must either express the extracellular receptor (α -component) or the cells may be exposed to the test agent in the presence of the soluble receptor component. Thus, for example, in an assay system designed to identify agonists or antagonists of CNTF, the cell may express the α - component CNTFR α , the β -components gp130 and LIFR β and a signal transducing component such as Jak1. The cell is exposed to test agents, and the tyrosine phosphorylation of either the β - components or the signal transducing component is

compared to the phosphorylation pattern produced in the presence of CNTF. Alternatively, the tyrosine phosphorylation which results from exposure to a test agent is compared to the phosphorylation which occurs in the absence of the test agent. Alternatively, an assay system, for example, for IL-6 may involve exposing a cell that expresses the β -component gp130 and a signal transducing protein such as Jak1, Jak2 or Tyk2 to a test agent in conjunction with the soluble IL-6 receptor.

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In another embodiment of the invention the above approaches are used to develop a method for screening for small molecule antagonists that act at 10 various steps in the process of ligand binding, receptor complex formation, and subsequent signal transduction. Molecules that potentially interfere with ligand-receptor interactions are screened by assessing interference of complex formation between the soluble receptors and ligand as described above. Alternatively, cell-based assays in which IL-6 or CNTF induce 15 response of a reporter gene are screened against libraries of small molecules or natural products to identify potential antagonists. Those molecules showing antagonist activity are rescreened on cell-based assays responding to other factors (such as GM-CSF or factors like Neurotrophin-3 that activate receptor tyrosine kinases) to evaluate their specificity against 20 the CNTF/IL-6/OSM/LIF family of factors. Such cell-based screens are used to identify antagonists that inhibit any of numerous targets in the signal transduction process.

In one such assay system, the specific target for antagonists is the interaction of the Jak/Tyk family of kinases [Firmbach-Kraft, Oncogene 5: 1329-1336 (1990); Wilks, et al., Mol. Cell. Biol. 11:2057-2065 (1991)] with the receptor β subunits. As described above, LIFRβ and gp130 preassociate with members of the Jak/Tyk family of cytoplasmic protein tyrosine kinases, which become activated in response to ligand-induced β component dimerization Stahl, et al. Science 263:92-95 (1993). Thus small molecules that could enter the cell cytoplasm and disrupt the interaction

between the \beta component and the Jak/Tyk kinase could potentially block all subsequent intracellular signaling. Such activity could be screened with an in vitro scheme that assessed the ability of small molecules to block the interaction between the relevant binding domains of purified β component and Jak/Tyk kinase. Alternatively, one could easily screen for molecules that could inhibit a yeast-based assay of β component binding to Jak/Tyk kinases using the two-hybrid interaction system [Chien, et al., Proc. Natl. Acad. Sci. 88: 9578-9582 (1991)]. In such a system, the interaction between two proteins (\beta component and Jak/Tyk kinase or relevant domains thereof in this example) induces production of a convenient marker such as β - galactosidase. Collections of small molecules are tested for their ability to disrupt the desired interaction without inhibiting the interaction between two control proteins. The advantage of this screen would be the requirement that the test compounds enter the cell before inhibiting the interaction between the β component and the Jak/Tyk kinase.

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The CNTF family antagonists described herein either bind to, or compete with the cytokines CNTF and IL-6. Accordingly, they are useful for treating diseases or disorders mediated by CNTF or IL-6. For example, therapeutic uses of IL-6 antagonists would include the following:

1) In osteoporosis, which can be exacerbated by lowering of estrogen levels in post-menopausal women or through ovariectomy, IL-6 appears to be a critical mediator of osteoclastogenesis, leading to bone resorption [Horowitz, Science 260: 626-627 (1993); Jilka, et al., Science 257: 88-91 (1992)]. Importantly, IL-6 only appears to play a major role in the estrogen-depleted state, and apparently is minimally involved in normal bone maintenance. Consistent with this, experimental evidence indicates that function-blocking antibodies to IL-6 can reduce the number of osteoclasts [Jilka, et al. Science 257: 88-91 (1992)]. While estrogen replacement therapy is also used, there appear to be side effects that may include increased risk of

endometrial and breast cancer. Thus, IL-6 antagonists as described herein would be more specific to reduce osteoclastogenesis to normal levels.

2) IL-6 appears to be directly involved in multiple myeloma by acting in either an autocrine or paracrine fashion to promote tumor formation [van Oers, et al., Ann Hematol. 66: 219-223 (1993)]. Furthermore, the elevated IL-6 levels create undesirable secondary effects such as bone resorption, hypercalcemia, and cachexia; in limited studies function-blocking antibodies to IL-6 or IL-6Ra have some efficacy [Klein, et al., Blood 78: 1198-1204 (1991); Suzuki, et al., Eur. J. Immunol. 22:1989-1993 (1992)]. Therefore, IL-6 antagonists as described herein would be beneficial for both the secondary effects as well as for inhibiting tumor growth.

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3) IL-6 may be a mediator of tumor necrosis factor (TNF) that leads to cachexia associated with AIDS and cancer [Strassmann, et al., J. Clin. Invest. 89: 1681-1684 (1992)], perhaps by reducing lipoprotein lipase activity in adipose tissue [Greenberg, et al., Cancer Research 52: 4113-4116 (1992)]. Accordingly, antagonists described herein would be useful in alleviating or reducing cachexia in such patients.

Effective doses useful for treating these or other CNTF family related diseases or disorders may be determined using methods known to one 20 skilled in the art [see, for example, Fingl, et al., The Pharmacological Basis of Therapeutics, Goodman and Gilman, eds. Macmillan Publishing Co., New York, pp. 1-46 ((1975)]. Pharmaceutical compositions for use according to the invention include the antagonists described above in a pharmacologically acceptable liquid, solid or semi-solid carrier, linked to a 25 carrier or targeting molecule (e.g., antibody, hormone, growth factor, etc.) and/or incorporated into liposomes, microcapsules, and controlled release preparation (including antagonist expressing cells) prior to administration in vivo. For example, the pharmaceutical composition may comprise one or more of the antagonists in an aqueous solution, such as sterile water, 30 saline, phosphate buffer or dextrose solution. Alternatively, the active agents may be comprised in a solid (e.g. wax) or semi-solid (e.g. gelatinous) formulation that may be implanted into a patient in need of such

treatment. The administration route may be any mode of administration known in the art, including but not limited to intravenously, intrathecally, subcutaneously, by injection into involved tissue, intraarterially, intranasally, orally, or via an implanted device.

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Administration may result in the distribution of the active agent of the invention throughout the body or in a localized area. For example, in some conditions which involve distant regions of the nervous system, intravenous or intrathecal administration of agent may be desirable. In some situations, an implant containing active agent may be placed in or near the lesioned area. Suitable implants include, but are not limited to, gelfoam, wax, or microparticle-based implants.

EXAMPLES

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EXAMPLE 1: CNTF COMPETES WITH IL-6 FOR BINDING TO GP130

MATERIALS AND METHODS

- Materials. A clone of PC12 cells that respond to IL-6 (PC12D) was obtained from DNAX. Rat CNTF was prepared as described [Masiakowski, et al., J. Neurochem. 57:1003-10012 (1991)]. IL-6 and sIL-6Rα were purchased from R & D Systems. Antisera was raised in rabbits against a peptide derived from a region near the C-terminus of gp130 (sequence:
- 25 CGTEGQVERFETVGME) [SEQ. ID. NO. 2] by the method described (Stahl, et al. J. Biol. Chem. 268:7628-7631 (1993). Anti-phosphotyrosine monoclonal 4G10 was purchased from UBI, and reagents for ECL from Amersham.
- Signal Transduction Assays. Plates (10 cm) of PC12D were starved in serum-free medium (RPMI 1640 + glutamine) for 1 hour, then incubated with IL-6 (50 ng/mL) + sIL-6R (1 mg/mL) in the presence or absence of

added rat CNTF at the indicated concentrations for 5 minutes at 37°C. Samples were then subjected to anti-gp130 immunoprecipitation, SDS PAGE, and anti-phosphotyrosine immunoblotting as described (Stahl, et al. J. Biol. Chem. 268:7628-7631 (1993).

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RESULTS

The ability of CNTF to block IL-6 responses was measured using a PC12 cell line (called PC12D) that expresses IL-6Rα, gp130, and CNTFRα, but not LIFRB. As one would predict, these cells respond to IL-6, but not to CNTF 10 (Fig. 2) since LIFRβ is a required component for CNTF signal transduction [Davis, et al., Science 260: 59-63 (1993)]. In accordance with results on other cell lines [Ip, et al., Cell 69: 1121-1132 (1992)], PC12D cells give tyrosine phosphorylation of gp130 (as well as a variety of other proteins called CLIPs) in response to 2 nM IL-6 (Fig. 2). Addition of recombinant soluble 15 IL-6Rα (sIL-6Rα) enhances the level of gp130 tyrosine phosphorylation, as has been reported in some other systems [(Taga, et al., Cell 58: 573-581 (1989)]. However, addition of 2 nM CNTF simultaneously with IL-6 severely diminishes the tyrosine phosphorylation of gp130. Although a slight gp130 phosphorylation response remains in the presence of CNTF, IL-6, and sIL-6Rα, it is eliminated if the CNTF concentration is increased fourfold to 8 nM. Thus, in IL-6 responsive cells that contain CNTFRα but no LIFRβ, CNTF is a rather potent antagonist of IL-6 action.

25 EXAMPLE 2. BINDING OF CNTF TO THE CNTFRα:β

MATERIALS AND METHODS

Scatchard Analysis of CNTF Binding. 125I-CNTF was prepared and purified as described [Stahl et al. JBC 268: 7628-7631 (1993)]. Saturation binding studies were carried out in PC12 cells, using concentrations of 125I-

CNTF ranging from 20pM to 10nM. Binding was performed directly on a monolayer of cells. Medium was removed from wells and cells were washed once with assay buffer consisting of phosphate buffered saline (PBS; pH 7.4), 0.1mM bacitracin, 1mM PMSF, 1mg/ml leupeptin, and 1mg/ml BSA. Cells were incubated in ¹²⁵I-CNTF for 2 hours at room temperature, followed by 2 quick washes with assay buffer. Cells were lysed with PBS containing 1% SDS and counted in a Packard Gamma Counter at 90-95% efficiency. Non-specific binding was defined by the presence of 100-fold excess of unlabelled CNTF. Specific binding ranged from 70% to 95%.

RESULTS

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The equilibrium constant for binding of CNTF to CNTFRα:β1 was estimated from Scatchard analysis of iodinated CNTF binding on PC12D 15 cells (Figure 3). The data is consistent with a 2 site fit having dissociation constants of 9 pM and 3.4 nM. The low affinity site corresponds to interaction of CNTF with CNTFR α , which has a Kd near 3 nM [(Panayotatos, et al., J. Biol. Chem. 268: 19000-19003 (1993)]. We interpret the high affinity complex as the intermediate containing CNTF, CNTFR α , 20 and gp130. A Ewing sarcoma cell line (EW-1) which does contain CNTFRα, gp130, and LIFRβ, and therefore gives robust tyrosine phosphorylation in response to CNTF, displays a very similar two site fit with dissociation constants of 1 nM and 10. Thus it is apparent that CNTF binds with equally high affinity to a complex containing only CNTFRa 25 and gp130, as it does to a complex which additionally contains LIFR\$\beta\$, thus demonstrating the feasibility of creating the $sR\alpha:\beta$ antagonists described herein.

EXAMPLE 3. METHODS OF PRODUCING CYTOKINE LIGAND TRAPS

Virus Stock Production

- 5 SF21 insect cells obtained from *Spodoptera frugiperda* were grown at 27°C in Gibco SF900 II medium to a density of 1x10⁶ cells/mL. The individual virus stock for either GP130-Fc-His6 (Figure 4) or IL6Ra-Fc (Figure 5) was added to the bioreactor to a low multiplicity 0.01-0.1 PFU/cell to begin the infection. The infection process was allowed to continue for 5-7 days allowing maximum virus replication without incurring substantial cell lysis. The cell suspension was aseptically aliquoted into sterile centrifuge bottles and the cells removed by centrifugation. The cell-free supernatant was collected in sterile bottles and stored at 4°C until further use.
- The virus titer was determined by plaque assay as described by O'Reilly, Miller and Luckow. The method is carried out in 60mm tissue-culture dishes which are seeded with 2x10⁶ cells. Serial dilutions of the virus stock are added to the attached cells and the mixture incubated with rocking to allow the virus to adsorb to individual cells. An agar overlay is added and plates incubated for 5 7 days at 27°C. Staining of viable cells with neutral red revealed circular plaques resulting which were counted to give the virus titer.

Coinfection of Cells for Protein Production

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Uninfected SF21 Cells were grown in a 60L ABEC bioreactor containing 40L of SF900 II medium. Temperature was controlled at 27°C and the dissolved oxygen level was maintained at 50% of saturation by controlling the flowrate of oxygen in the inlet gas stream. When a density of 2x106 cells/mL was reached, the cells were concentrated within the bioreactor to a volume of 20L using a low shear steam sterilizable pump with a tangential flow filtration device with Millipore Prostak 0.65 micron

membranes. After concentration fresh sterile growth medium is slowly added to the bioreactor while the filtration system continues to remove the spent growth medium by diafiltration. After two volume exchanges (40L) have been carried out an additional 20L of fresh medium was added to the bioreactor to resuspend the cells to the original volume of 40L. The cell density was determined once again by counting viable cells using a hemacytometer.

The required amount of each virus stock was calculated based on the cell
density, virus titer and the desired multiplicity of infection (MOI). Virus
stock ratios of 5:1, 5:2, 10:2 and 10:4, IL6Rα-Fc to GP130-Fc-His6 all resulted
in production of significant amounts of heterodimer. The ideal virus
stock ratio is highly dependent on the ease of purification of the
heterodimer from each of the two homodimers. The IL6Rα-Fc

15 homodimer is relatively easy to remove downstream by immobilized
metal affinity chromatography. Virus infection ratios have been chosen to
minimize the formation of the GP130-Fc-His6 homodimer which is more
difficult to clear downstream. The relative amount of GP130-Fc-His6 virus
stock chosen for infection has increased with successive batches as the

20 purification method for clearing the resultant homodimer has improved.

The virus stocks were aseptically mixed in a single vessel then transferred to the bioreactor. This results in synchronous infection of the SF21 cells. The infection is allowed to proceed for three to four days, allowing sufficient time for maximal production of the heterodimer protein.

Recovery and Protein A Chromatographic Purification

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At the conclusion of the infection phase of the bioreactor process the cells were concentrated in the bioreactor using a 10 ft² Millipore Prostak filter (0.65 micron) pore size. The cell-free permeate passing through the filter was collected in a clean process vessel. At the conclusion of the filtration

operation the pH of permeate stream, containing the protein product, was adjusted to 8.0 with 10N NaOH. The resultant precipitate was removed by forcing the extract through a 0.8 micron depth filter (Sartorious), followed by a 0.2 micron filter. Sufficient 0.5M EDTA stock was added to give a final concentration of 5mM. The filtered protein solution was loaded onto a 10 cm diameter column containing 100-200 mL of Pharmacia Protein A Sepharose 4 Fast Flow, equilibrated with PBS. Protein A has a very high affinity for the Fc-Fc domain of each of the 3 recombinant protein products, allowing them to bind while other proteins in the cell-free extract flow through the column. After loading the column was washed to baseline with PBS containing an additional 350mM NaCl. The IgG-Fc tagged proteins were eluted at low pH, either with 0.5M acetic acid or with a decreasing pH gradient of 0.1M citric acid and 0.2M disodium phosphate buffers. Tris base or disodium phosphate was added to the eluted protein to avoid prolonged exposure to low pH conditions.

The pooled protein was diafiltered into PBS or HEPES buffer and derivitized with 1 mM iodoacetamide to protect the exposed sulfhydryl group on the free cysteine near the hinge region of each Fc domain. This prevents disulfide mediated aggregation of proteins. A 6 ft² Millipore spiral wound ultrafiltration membrane with nominal 30 kiloDalton cutoff was used to perform the buffer exchange. The total protein was determined by UV absorbance at 280 nm using the diafiltration buffer as a blank. The relative amounts of heterodimer and two homodimer proteins were determined by SDS PAGE gel electrophoresis using a 6% Tris-Glycine gel (Novex). Gels were Coomassie-stained then transferred into destain solution overnight. A Shimadzu scanning densitometer was used to determine the relative intensity of the individual protein bands on the SDS PAGE gel. The peak area ratios are used to compute the fraction of heterodimer and each of the homodimers in the column pool fractions.

Immobilized Metal Affinity Chromatographic Purification

The six histidine residues on the C-terminus of the GP130-Fc-His6 fusion protein provides an excellent molecular handle for separation of the heterodimeric IL6 antagonist from the two homodimers. The imidazole 5 group on each of the C-terminal histidines of the GP130-Fc-His6 moiety has a strong binding constant with several divalent metals, including copper, nickel, zinc, cobalt, iron and calcium. Since the IL6Rα-Fc homodimer has no C-terminal histidine residues, it clearly has the lowest affinity. The IL6Rα-Fc-GP130-Fc-His6 heterodimer has a single stand set 10 six histidines giving it greater affinity for the metal, while the GP130-Fc-His6 homodimer has two sets of six histidines each giving it the highest affinity of the three IgG tagged proteins to the metal affinity column. Selective elution of the three proteins with increasing amounts of imidazole in the elution buffer therefore elutes the proteins in the 15 following order:

- 1. IL6Rα-Fc homodimer
- 2. IL6Rα-Fc-GP130-Fc-His heterodimer
- 3. GP130-Fc-His homodimer

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A 26 mm diameter column containing 100 mL of Pharmacia Chelating Sepharose Fast Flow was saturated with a solution of nickel sulfate until a significant green color is observed in the column eluate. The column is then washed with several column volumes of deionized water, then equilibrated with 50 mM HEPES, 40mM imidazole, pH 8.0. The binding of imidazole to the immobilized nickel results in a green to blue color change. Imidazole was added to the protein load to a final concentration of 40mM. Addition of imidazole to the protein load reduces the binding of IL6Rα-Fc homodimer, increasing the surface area available for the remaining two species. After loading, the column was washed with

several column volumes of 50 mM HEPES, 80mM imidazole, pH 8.0 until a steady baseline was reestablished. The heterodimer was selectively eluted with 50 mM HEPES, 150mM imidazole, pH 8.0 over several column volumes. The protein fractions were pooled and diafiltered into PBS as described in the section above.

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EXAMPLE 4. ALTERNATIVE METHODS OF CONSTRUCTING LIGAND TRAPS

- As described above, receptor activation by CNTF, and analogously by IL-6 10 and IL-11, follows an ordered sequence of binding events (Figure 6). The cytokine initially binds to its cognate R α with low affinity (Kd = 3 to 10 nM); this is a required step - cells which do not express the cognate Rα do not respond to the cognate cytokine. The cytokine•Rα complex associates 15 with the first signal transducing component, gp130, to form a high affinity complex (Kd in the order of 10 pM for the CNTF•CNTFRa•gp130 complex). This complex does not transduce signal, as it is the dimerization of the signal transducing components that brings about signaling (Stahl and Yancopoulos, J. Neurobiology 25: 1454-1466 (1994); Stahl et al., Science 267:1349-1353 (1995); Davis et al., Science 260:1805-1808 (1993); Stahl et al., 20 Science 263:92-95 (1994); Murakami, et al. Science 260:1808-1810 (1993). At least in the case of IL-6, the cytokine • Ra • signal transducer heterotrimeric complex subsequently associates with another like complex, to form a hexameric complex (Figure 6) (Ward et al., J. Biol. Chem. 269:23286-23289 (1994). The resulting dimerization of the signal transducers - gp130 in the 25 case of IL-6 (Murakami et al., Science 260:1808-1810 (1993) and IL-11, gp130 and LIFR in the case of CNTF (Davis et al., Science 260:1805-1808 (1993) brings about signal transduction.
- The initial heterodimeric molecules made comprised a soluble Rαcomponent linked to the extracellular domain of gp130. These molecules

were shown to mimic the high affinity cytokine •R α •gp130 complex and behave as a high affinity antagonist of their cognate cytokine (Figure 7). To make these molecules, the extracellular domain of gp130 was paired with the extracellular domain of the α -receptor components for IL-6 and CNTF,

- IL-6Rα and CNTFRα respectively. To link the Rα with the extracellular domain of gp130, the soluble Rα-components and gp130 were fused to the Fc portion of human IgG1 to produce Rα-Fc and gp130-Fc respectively. The Fc domain was chosen primarily but not solely because it naturally forms disulfide-linked dimers. Heterodimeric molecules comprising Rα-
- Fc•gp130-Fc were expressed, purified and shown to behave as highly potent antagonists of their cognate ligand. Furthermore, these molecules were found to be highly specific for their cognate cytokine since it is the choice of the α-receptor component which specifies which cytokine is bound and trapped (there is no measurable binding of the cytokine to gp130 in the absence of the appropriate Rα).

Here we describe an extension of this technology which allows the engineering of different heteromeric soluble receptor ligand traps which by virtue of their design may have additional beneficial characteristics such as stability, Fc-receptor-mediated clearance, or reduced effector functions (such as complement fixation). Furthermore, the technology described should prove suitable for the engineering of any heteromeric protein in mammalian or other suitable protein expression systems, including but not limited to heteromeric molecules which employ receptors, ligands, and catalytic components such as enzymes or catalytic antibodies.

MATERIALS AND METHODS

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Genetic engineering of heteromeric immunoglobulin heavy/light chain soluble receptor-based ligand traps for IL-6.

The IL-6 traps described here were engineered using human gp130, human IL-6 α -receptor (IL-6R α), the constant region of the heavy chains (C γ) of human IgG1 (Cγ1) (Lewis et al., Journal of Immunology 151:2829-2838 (1993) or IgG4 (Cy4) with or without a join-region (J), and the constant regions of kappa (κ) and lambda (λ) (Cheung, et al., Journal of Virology 66:6714-6720 (1992) light chains of human immunoglobulin (Ig), also with or without a different j-peptide (j). This design takes advantage of the natural ability of the Cy domain to heterodimerize with κ or λ light chains. The heterodimerization of Cy with the light chain occurs between the CH1 domain of Cy and the constant region of the light chain (CL), and is stabilized by covalent linking of the two domains via a single disulfide bridge. We reasoned that, like the Fc domain of human IgG1, the combination of Cy with CL could be used to produce disulfide linked heteromeric proteins comprised of the extracellular domain of gp130 on one chain and the extracellular domain of IL-6R α on the other chain. Like their Fc-based counterparts, such proteins were postulated to be high affinity ligand traps for IL-6 and as a result to inhibit the interaction of IL-6 with the native receptor on IL-6-responsive cells, thus functioning as IL-6 antagonists. Furthermore, constructs employing the full length Cy region would, much like antibodies, form homodimers of the Cy chain, giving rise to antibody-like molecules comprising of two "light chains" and two "heavy chains" (Figure 8). The potential advantage of this design is that it may more closely mimic the IL-6•IL-6Rα•gp130 complex and may display a higher affinity for the ligand than comparable single heterodimers. An additional design is incorporated by using truncated versions of Cy, comprised only of the CH1 domain. These will form heterodimeric molecules with receptor-κ fusion proteins, and will thus resemble the Fab fragment of antibodies.

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All the soluble receptor-Ig chimeric genes may be engineered in plasmid vectors including, but not limited to, vectors suitable for mammalian expression (COS monkey kidney cells, Chinese Hamster Ovary cells [CHO], and ras-transformed fibroblasts [MG-ras]) and include a Kozak sequence (CGC CGC CAC CAT GGT G) at the beginning of each chimeric gene for efficient translation. Engineering was performed using standard genetic engineering methodology. Each construct was verified by DNA sequencing, mammalian expression followed by western blotting with suitable antibodies, biophysical assays that determine ligand binding and dissociation, and by growth inhibition assays (XG-1, as described later). Since the domains utilized to engineer these chimeric proteins are flanked by appropriate restriction sites, it is possible to use these domains to engineer other chimeric proteins, including chimeras employing the extracellular domains of the receptors for factors such as IL-1, IL-2, IL-3, IL-4, IL-5, GM-CSF, LIF, IL-11, IL-15, IFNγ, TGFβ, and others. The amino acid coordinates for each component utilized in making the IL-6 traps are listed below (Note: numbering starts with the initiating methionine as #1; long sequences are listed using the single letter code for the twenty amino acids):

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(a) Constructs employing human gp130:

- (i) gp130-Cγ1 was engineered by fusing in frame the extracellular domain of gp130 (amino acids 1 to 619) to a Ser-Gly bridge, followed by the 330 amino acids which comprise Cγ1 and a termination codon (Figure 9).
- (ii) gp130-J-Cγ1 was engineered in the same manner as gp130-Cγ1 except that a J-peptide (amino acid sequence: GQGTLVTVSS) was inserted between the Ser-Gly bridge and the sequence of Cγ1 (see Figure 9).
 - (iii) gp130Δ3fibro-Cγ1 was engineered by fusing in frame the extracellular domain of gp130 without its three fibronectin-like domains (Figure 10).
- The remaining part of this chimeric protein is identical to gp130-Cγ1.

(iv) gp130-J-CH1 was engineered in a manner identical for that described for gp130-Cγ1, except that in place of the Cγ1 region only the CH1 part of Cγ1 has been used (Figure 11). The C-terminal domain of this construct includes the part of the hinge that contains the cysteine residue

- responsible for heterodimerization of the heavy chain of IgG with a light chain. The part of the hinge that contains the two cysteines involved in Cγ1 homodimerization has been deleted along with the CH2 and CH3 domains.
- (v) gp130-Cγ4 was engineered in a manner identical to that described for gp130-Cγ1, except that Cγ4 was used in place of Cγ1 (Figure 12). In addition, an RsrII DNA restriction site was engineered at the hinge region of the Cγ4 domain by introducing two silent base mutations. The RsrsII site allows for other desired genetic engineering manipulations, such as the construction of the CH1 equivalent of gp130-Cγ4.
- (vi) gp130- κ was engineered in a manner identical to that described for gp130-Cγ1, except that the constant region of the κ light chain of human Ig was used in place of Cγ1 (Figure 13).
 - (vi) gp130-J- κ was engineered in a manner identical to that described for gp130-J- κ , except that a j-peptide (amino acid sequence: TFGQGTKVEIK) was inserted between the Ser-Gly bridge and the κ -region.
 - (viii) gp130- λ was engineered in a manner identical to that described for gp130-C γ 1, except that the constant region of the λ light chain (Cheung, et al., Journal of Virology 66:6714-6720 (1992) of human Ig was used in place of C γ 1 (Figure 14).

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- (b) Constructs employing human IL-6Rα:
- (i) IL6R α -C γ 1 was engineered by fusing in frame amino acids 1 to 358 of IL-6R α (Yamasaki et al., Science 241:825-828 (1988), which comprise the

extracellular domain of IL-6R α (Figure 15), to an Ala-Gly bridge, followed by the 330 amino acids which comprise C γ 1 and a termination codon.

- (ii) IL6R α - κ was engineered as described for IL6R α -C γ 1, except that the κ -domain (Figure 13) utilized for gp130- κ was used in place of C γ 1.
- 5 (iii) IL6Rα-j- κ was engineered as described for IL6Rα- κ except that the j-peptide described for gp130-j- κ was placed between the Ala-Gly bridge and the κ -domain.
 - (iv) Three additional constructs, IL6Rα313-Cγ1, IL6Rα313-κ, and IL6Rα313-j-κ, were engineered as using a truncated form of IL-6Rα comprised of amino acids 1 to 313 (Figure 16). Each of these constructs were made by fusing in frame IL6Rα313 with a Thr-Gly bridge followed by the Cγ1, κ-, and j-κ-domains described above. These constructs were engineered in order to complement the gp130Δ3fibro-derived constructs.

15 Expression and purification of ligand traps

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To produce covalently linked heterodimers of soluble gp130 and soluble IL-6Rα, gp130-Ig chimeric proteins were co-expressed with appropriate IL-6Rα-Ig chimeric proteins in complementing pairs. Co-expression was achieved by co-transfecting the corresponding expression vectors into suitable mammalian cell lines, either stably or transiently. The resulting disulfide-linked heterodimers were purified from conditioned media by several different methods, including but not limited to affinity chromatography on immobilized Protein A or Protein G, ligand-based affinity chromatography, ion exchange, and gel filtration.

An example of the type of methods used for purification of a heavy/light receptor fusion protein is as follows: gp130-C γ 1•IL-6R α - κ was expressed in COS cells by co-transfecting two different vectors, encoding gp130-C γ 1 and

IL-6R α - κ respectively. Serum-free conditioned media (400 ml) were collected two days post-transfection and C γ 1-bearing proteins were purified by affinity chromatography over a 1ml Protein A Sepharose (Pharmacia). The material generated in this step was further purified by a second affinity chromatography step over a 1 ml NHS-activated Sepharose (Pharmacia) which was derivatized with recombinant human IL-6, in order to remove gp130-C γ 1 dimer from gp130-C γ 1 •IL-6R α - κ complexes (the gp130-C γ 1 dimer does not bind IL-6). Proteins generated by this method were more than 90% pure, as evidenced by SDS-PAGE followed by silverstaining (Figure 17). Similar protocols have been employed successfully towards the purification of other heavy/light receptor heterodimers.

RESULTS

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Biological activity of immunoglobulin heavy/light chain receptor fusion antagonists

The purified ligand traps were tested for their ability to bind IL-6 in a variety of different assays. For example, the dissociation rate of IL-6 bound to the ligand trap was measured in parallel with the dissociation rate of IL-20 6 from the anti-IL-6 monoclonal neutralizing antibody B-E8 [Brochier, et al., Int. J. Immunopharmacology 17:41-48 (1995), and references within]. An example of this type of experiment is shown in Figure 18. In this experiment 20 pM ¹²⁵I-IL-6 (1000 µCi/mmol; Amersham) was preincubated with 500 pM of either gp130-Cγ1 • IL-6Rα-κ or mAb B-E8 for 20 25 hours. At this point a 1000-fold excess (20 nM) of "cold" IL-6 was added. Periodically, aliquots of the reaction were removed, the ligand trap or B-E8 were precipitated with Protein G-Sepharose, and the number of cpm of 125I-IL-6 that remained bound was determined. Clearly, the dissociation rate of human ¹²⁵I-IL6 from the ligand trap was very slow - after three 30 days, approximately 75% of the initial counts were still bound to the ligand

trap. In contrast, less than 5% of the counts remained associated with the antibody after three days. This result demonstrates that the dissociation rate of the ligand from these ligand traps is very slow.

In a different set of experiments the ability of the ligand traps to multimerize in the presence of ligand was tested. An example of this is shown in Figure 19. IL-6-induced association of gp130-Fc•IL-6Rα-Fc with gp130-CH1•IL-6Rα-κ was determined by testing whether gp130-CH1•IL-6Rα-κ, which does not by itself bind Protein A, could be precipitated by Protein A-Sepharose in the presence of gp130-Fc•IL-6Rα-Fc in an IL-6-10 depended manner (Figure 9). Precipitation of gp130-CH1•IL-6Rα-κ by Protein A-Sepharose was determined by western blotting with an antikappa specific HRP conjugate, which does not detect gp130-Fc•IL-6Rα-Fc. gp130-CH1•IL-6Rα-κ could be precipitated by Protein A-Sepharose only when both gp130-Fc•IL-6Rα-Fc and IL-6 were present. This result 15 conclusively indicates that IL-6 can induce ligand trap multimerization, and further indicate that the ligand trap can mimic the hexameric cytokine • Ra • signal transducer complex (Figure 1). Ligand-induced multimerization may play a significant role in the clearance of cytokine • ligand trap complexes in vivo. 20

The biological activity of the different ligand traps may be further tested in assays which measure ligand-depended cell proliferation. Several cell proliferation assays exist for IL-6 and they employ cell lines such as B9, CESS, or XG-1. An example of this type of assay using the XG-1 cell line is presented below: XG-1 is a cell line derived from a human multiple myeloma (Zhang, et al., Blood 83:3654-3663 (1994). XG-1 depends on exogenously supplied human IL-6 for survival and proliferation. The EC50 of IL-6 for the XG-1 line is approximately 50 pmoles/ml. The ability of several different IL-6 traps to block IL-6-depended proliferation of XG-1

cells was tested by incubating increasing amounts of purified ligand traps with 50 pg/ml IL-6 in XG-1 cultures. The ligand traps which were tested had been expressed and purified by methods similar to those described above. All of the ligand traps tested were found to inhibit IL-6-dependent proliferation of XG-1 in a dose dependent manner (Figure 20). Of the five different traps tested gp130-Cγ1•IL-6Rα-κ was the most active and essentially display the same neutralizing activity towards IL-6 as the antibody B-E8. As little as a 10-fold molar excess of either gp130-Cy1 • IL-6Rα-κ or B-E8 completely blocked the activity of IL- 6 (a reading of A570-650 = 0.3 AU corresponds to no proliferation of the XG-1 cells). At a 100fold molar excess all of the ligand traps tested completely blocked the activity of IL-6. This observed inhibition is highly selective as neither a gp130-Fc•CNTFRα-Fc ligand trap which blocks CNTF activity, nor gp130-Fc homodimer exhibit any blocking activity towards IL-6 even when used at a 1000-fold molar excess over IL-6 (data not shown). This data demonstrates that the heteromeric immunoglobulin heavy/light chain receptor-based ligand traps function as selective high affinity antagonists of their cognate ligand.

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20 EXAMPLE 5 - CLONING OF FUSION POLYPEPTIDE COMPONENTS

The extracellular domains of the human cytokine receptors were obtained by standard PCR techniques using tissue cDNAs (CLONTECH), cloned into the expression vector, pMT21 (Genetics Institute, Inc.), and the sequences were sequenced by standard techniques using an ABI 373A DNA sequencer and Taq Dideoxy Terminator Cycle Sequencing Kit (Applied Biosystems, Inc., Foster City, CA). For the IL-4Rα, nucleotides 241 through 868 (corresponding to the amino acids 24-231) from the Genbank sequence, X52425, were cloned. For the IL-2Rγ, nucleotides 15 through 776 (corresponding to amino acids 1-233) from the Genbank sequence, D11086, were cloned. For the IL-6Rα, nucleotides 52 through 1044 (corresponding

to the amino acids 1-331) from the Genbank sequence, X52425, were cloned. For gp130, nucleotides 322 through 2112 (corresponding to the amino acids 30-619) from the Genbank sequence, M57230, were cloned. For the IL-1RAcP, nucleotides 1 through 1074 (corresponding to the amino acids 1-358) from the Genbank sequence, AB006357, were cloned. For the IL-1RI, nucleotides 55 through 999 (corresponding to the amino acids 19-333) from the Genbank sequence, X16896, were cloned.

EXAMPLE 6 - PRODUCTION OF FUSION POLYPEPTIDES (CYTOKINE 10 TRAPS)

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The nucleotide sequences encoding the cytokine traps were constructed from the individual cloned DNAs (described *supra*) by standard cloning and PCR techniques. In each case, the sequences were constructed in frame such that the sequence encoding the first fusion polypeptide component was fused to the sequence encoding the second fusion polypeptide component followed by an Fc domain (hinge, CH2 and CH3 region of human IgG1) as the multimerizing component. In some cases extra nucleotides were inserted in frame between sequences encoding the first and second fusion polypeptide components to add a linker region between the two components (See Figure 21A - Figure 21D - trap 424; Figure 24A - Figure 24F - trap 412; and Figure 26A - Figure 26E - trap 569).

For the IL-4 traps, 424 (Figure 21A - Figure 21D), 603 (Figure 22A - Figure 22D) and 622 (Figure 23A - Figure 23D), the IL-2Rγ component is 5′, followed by the IL4Rα component and then the Fc component. For the IL-6 traps, 412 (Figure 24A - Figure 24F) and 616 (Figure 25A - Figure 25F), the IL-6Rα component is 5′ followed by the gp130 component and then the Fc domain. For the IL-1 trap 569 (Figure 26A - Figure 26E), the IL-1RAcP component is 5′ followed by the IL-1RI component and then the Fc domain. The final constructs were cloned into the mammalian expression vector pCDNA3.1 (STRATAGENE).

In the 569 sequence (Figure 26A - Figure 26E), nucleotides 1-1074 encode the IL1RAcP component, nucleotides 1075 -1098 encode a linker region, nucleotides 1099-2043 encode the IL1RI component and nucleotides 2044-2730 encode the Fc domain.

In the 412 sequence (Figure 24A - Figure 24F), nucleotides 1-993 encode the IL6Rα component, nucleotides 994-1023 encode a linker region, nucleotides 1024-2814 encode the gp130 component and nucleotides 2815-3504 encode the Fc domain.

In the 616 sequence (Figure 25A - Figure 25F), nucleotides 1-993 encode the IL6Rα component, nucleotides 994-2784 encode the gp130 component and nucleotides 2785-3474 encode the Fc domain.

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In the 424 (Figure 21A - Figure 21D) and 622 (Figure 23A - Figure 23D) sequences, nucleotides 1-762 encode the IL2R γ component, nucleotides 763-771 encode a linker region, nucleotides 772-1395 encode the IL4R α component and nucleotides 1396-2082 encode the Fc domain.

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Finally, in the 603 sequence (Figure 22A - Figure 22D), nucleotides 1-762 encode the IL2Ry component, nucleotides 763-1386 encode the IL4R α component and nucleotides 1387-2073 encode the Fc domain.

DNA constructs were either transiently transfected into COS cells or stably transfected into CHO cells by standard techniques well known to one of skill in the art. Supernatants were collected and purified by Protein A affinity chromatography and size exclusion chromatography by standard techniques. (See for example Harlow and Lane, Antibodies - A Laboratory Manual, Cold Spring Harbor Laboratory, 1988).

EXAMPLE 7: IL-4 BIOASSAY PROTOCOL USING TF-1 (ATCC) CELLS.

Reagents and Equipment Needed

5 MTT Dye Solution:

MTT(3-[4,5-Dimethylthiazole-2-yl]) (Sigma catalog# M2128) Working concentration: Dissolve 5 mg of anhydrous MTT in 200 ml PBS without Ca+2, Mg+2.

10 Sterile filter and store aliquoted at -20°C

Solubilization Solution:

For 1000 ml, combine 100 g SDS, 950 ml d H_20 , 50 ml Dimethyl Formamide, and 850 μ l concentrated HCl. Filter sterilize with a 0.45 μ m filter unit. Store at room temperature

TF-1 cell Growth Medium:

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RPMI 1640, 10% FBS, Pen/Strep, 2mM L-glutamine

Other:

0.4% Trypan Blue Stain, sterile tubes for dilutions, sterile 96 well cell culture plates (Falcon #3072), hemacytometer, centrifuge, ELISA plate reader, multichannel pipet for 15, 25, 50 and 100μl volume, sterile reagent reservoirs, sterile pipet tips, gloves.

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Assay Protocol

A. Preparation of Assay plates

Prepare sterile 96 well tissue culture plates to contain 50μl of growth medium per well with various concentrations of IL-4 and 10nM IL-4 antagonist. This can be done by preparing a working dilution of IL-4 that is 4 times the highest concentration to be assayed. In separate tubes, do a two-fold serial dilution of the IL-4. Add 25μl of each dilution to one row across the plate (i.e. row A gets highest concentration, row G gets lowest concentration). Add 25μl of growth medium without IL-4 to row H. Prepare the antagonists to be tested by making a stock that is 4 times the final concentration. Add 25μl to a triplicate set of IL-4 containing wells (columns 1,2,3, A through H). Be sure to include antagonist in row H.

2. As a positive control, leave one set with no antagonist. These wells will contain IL-4 and media only.

3. Incubate the plate for 1-2 hours at 37°C in a humidified 5% CO₂ incubator before preparing cells to be used for assay.

B. Preparation of Cells

- 4. Wash cells twice by centrifugation in assay medium free of growth factor.
 - 5. Determine cell number and trypan blue viability and suspend cells to a final concentration of 8×10^5 /ml in assay medium.
- 6. Dispense 50μl of the cell suspension (40,000 cells) into all wells of the plates. Total volume should now be 100μl/well.

7. Incubate the plate at 37°C for 68 hours in a humidified 5% CO₂ incubator.

C. Color Development

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- 8. After incubating for 68 hours, add $15\mu l$ of the MTT dye solution to each well.
- 9. Incubate the plate at 37°C for 4 hours in a humidified 5% CO₂ incubator.

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- 10. After 4 hours, add 100µl of the solubilization solution to each well. Allow the plate to stand overnight in a sealed container to completely solubilize the formazan crystals.
- 15 11. Record the absorbance at 570/650nm.

RESULTS

Figure 27 shows that an IL-4 trap designated 4SC375, which is a fusion polypeptide of IL-2R γ -scb-IL4R α -Fc Δ C1, is several orders of magnitude better as an IL-4 antagonist than IL4R α Fc Δ C1 alone in the TF1 cell bioassay.

Figure 28 shows that the IL-4 trap designated 4SC375 shows antagonistic activity in the TF1 cell bioassay equivalent to an IL-4 trap designated 4SC424 which is a fusion polypeptide of IL-2R γ -IL4R α -Fc Δ C1 having the IL-2R γ component flush with the IL-4R α component.

EXAMPLE 8: IL-6 BIOASSAY PROTOCOL USING XG-1 CELLS

30 Reagents and Equipment Needed

MTT Dye Solution:

MTT(3-[4,5-Dimethylthiazole-2-yl]) (Sigma catalog# M2128) Working concentration: Dissolve 5 mg of anhydrous MTT in 200 ml PBS without Ca^{+2} , Mg^{+2} .

Sterile filter and store aliquoted at -20°C

Solubilization Solution:

Filter sterilize with at 0.45 μm filter unit

Filter sterilize with at 0.45µm filter unit.

Store at room temperature

15 Assay Medium:

RPMI 1640, 10%FBS, Pen/Strep, 2mM L-glutamine, 50µM mercapto-ethanol.

20 Other:

0.4% Trypan Blue Stain, sterile tubes for dilutions, sterile 96 well cell culture plates (Falcon#3072), hemacytometer, centrifuge, ELISA plate reader, multichannel pipet for 15, 25, 50 and 100µl volume, sterile reagent reservoirs, sterile pipet tips, gloves.

Assay Protocol

A. Preparation of Assay plates

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1. Prepare sterile 96 well tissue culture plates to contain 50µl of growth medium per well with various concentrations of IL-6 and 10nM IL-6 antagonist. This can be done by preparing a working dilution of IL-6 that is

4 times the highest concentration to be assayed. In separate tubes, do a two-fold serial dilution of the IL-6. Add 25µl of each dilution to one row across the plate (i.e. row A gets highest concentration, row G gets lowest concentration). Add 25µl of growth medium without IL-6 to row H.

- Prepare the antagonists to be tested by making a stock that is 4 times the final concentration. Add 25µl to a triplicate set of IL-6 containing wells (columns 1,2,3, A through H). Be sure to include antagonist in row H. A typical IL-6 titration starts at 200ng/ml down to 3.1ng/ml.
- 2. As a positive control, leave one set with no antagonist. These wells contain IL-6 and media in place of antagonist.
 - 3. Incubate the plate 1-2 hours at 37oC in a humidified 5% CO₂ incubator before preparing cells to be used for assay.

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B. Preparation of Cells

4. Wash cells twice by centrifugation (5 min at 1000RPM) in assay medium free of growth factor.

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- 5. Determine cell number and trypan blue viability and suspend cells to a final concentration of $8 \times 10^5/\text{ml}$ in assay medium.
- 6. Dispense 50µl of the cell suspension (40000 cells) into all wells of the plates. Total volume should now be 100µl/well.
 - 7. Incubate the plate at 37°C for 68 hours in a humidified 5% CO₂ incubator.

30 <u>C. Color Development</u>

8. At 68 hours add 15µl of the dye solution to each well.

9. Incubate the plate at 37°C for 4 hours in a humidified 5% CO₂ incubator.

10. After 4 hours, add 100µl of the solubilization solution to each well. Allow the plate to stand overnight in a sealed container to completely solubilize the formazan crystals.

11. Record the absorbance at 570/650nm.

RESULTS

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Figure 29 shows that the IL6 trap (6SC412 IL6R-scb-gpx-FcΔC1) described in Figure 24A - Figure 24F is a better antagonist of IL-6 in the XG1 bioassay than the neutralizing monoclonal antibody to human IL-6 - BE8.

15 EXAMPLE 9: MRC5 BIOASSAY FOR IL1 TRAPS

MRC5 human lung fibroblast cells respond to IL-1 by secreting IL-6 and thus were utilized to assay the ability of IL-1 traps to block the IL-1-dependent production of IL-6. IL1 Trap 1SC569 (Figure 26A - Figure 26E) was tested against IL-1-RI.Fc which is the extracellular domain of the IL-1 Type I receptor fused to an Fc domain.

MRC5 cells are suspended at 1×10^5 cells per ml in medium and 0.1 ml of cells are plated (10,000 cells per well) into the wells of a 96 well tissue culture plate. Plates are incubated for 24 hours at 37°C in a humidified 5% CO_2 incubator.

IL-1 trap and recombinant human IL-1 at varying doses are pre-incubated in a 96 well tissue culture dish and incubated for 2 hours at 37°C. 0.1 ml of this mixture is then added to the 96 well plate containing the MRC5 cells such that the final concentration of IL-1 Trap is 10nM and the final

concentrations of the IL-1 ranges from 2.4 pM to 5nM. Control wells contain trap alone or nothing.

Plates are then incubated at 37°C for 24 hours in a humidified 5% CO₂ incubator. Supernatant is collected and assayed for levels of IL-6 using R&D Systems Quantikine Immunoassay Kit according to the manufacturer's instructions.

RESULTS

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Figure 30 shows that the trap 569 (Figure 26A - Figure 26E) is able to antagonize the effects of IL-1 and block the IL-6 production from MRC 5 cells upon treatment with IL-1. At a concentration of 10nM, the trap 569 is able to block the production of IL-6 up to an IL-1 concentration of 3nM. In contrast, the IL-1RI.Fc is a much poorer antagonist of IL-1. It is only able to block the effects of IL-1 up to about 10-20 pM. Thus, the trap 569 is approximately 100x better at blocking IL-1 than IL1RI.Fc.

EXAMPLE 10 - CONSTRUCTION OF IL-13/IL-4 SINGLE CHAIN TRAPS

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1. To create the IL-13/IL-4 dual trap designated IL-4R α .IL-13R α 1.Fc, the human IL-4R α extracellular domain (corresponding to nucleotides #1-693 of Figure 31A - Figure 31G) and the human IL-13R α 1 extracellular domain (corresponding to nucleotides #700-1665 of Figure 31A - Figure 31G) were amplified by standard PCR techniques and ligated into an expression vector pMT21 which contained the human Fc sequence (corresponding to nucleotides #1671-2355 of Figure 31A - Figure 31G), thus creating a fusion protein consisting of the IL-4R α , IL-13R α 1, and the hinge, CH2 and CH3 region of human IgG1 from the N to C terminus. In addition, a two amino acid linker (corresponding to nucleotides #694-699 of Figure 31A - Figure 31G) with the amino acid sequence SerGly was constructed in frame

between the IL-4Rα and the IL-13Rα1 and a two amino acid linker (corresponding to nucleotides #1666-1671 of Figure 31A - Figure 31G) with the amino acid sequence ThrGly was constructed in frame between the IL-13Rα1 and the Fc portion. All sequences were sequence-verified by standard techniques. The IL-4Rα.IL-13Rα1.Fc coding sequence was then subcloned into the expression vector pCDNA3.1 (Stratagene) using standard molecular biology techniques.

2. To create the IL-13/IL-4 dual trap designated IL-13Rα1.IL-4Rα.Fc, the IL-13Rα1 extracellular domain (corresponding to nucleotides #1-1029 of 10 Figure 32A - Figure 32G) and the human IL-4Rα (corresponding to nucleotides # 1060-1692 of Figure 32A - Figure 32G) were amplified by standard PCR techniques and ligated into the expression vector pJFE14, which contains the human Fc sequence (corresponding to nucleotides #1699-2382 of Figure 32A - Figure 32G) to create a fusion protein consisting 15 of the IL-13Rα1, IL-4Rα, and the hinge, CH2 and CH3 region of human IgG1 from the N to C terminus. In addition, a ten amino acid linker with the amino acid sequence GlyAlaProSerGlyGlyGlyArgPro (corresponding to nucleotide #1030-1059 of Figure 32A - Figure 32G) was constructed in frame between the IL-13Ral and the IL-4Ra and a two 20 amino acid linker (corresponding to nucleotides #1693-1698 of Figure 32A -Figure 32G) with the amino acid sequence SerGly was constructed in frame between IL-4Ra and the Fc portion. All sequences were sequence-verified using standard techniques. The coding sequence of IL-13Rα1.IL-4Rα.Fc was then subcloned into the expression vector pCDNA3.1 (Stratagene) 25 using standard molecular biology techniques.

EXAMPLE 11: EXPRESSION OF IL-4Rα.IL-13Rα1.Fc AND IL-13Rα1.IL-4Rα.Fc

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Large scale (1L) cultures of the pCAE801 (the DNA vector construct encoding IL-4Rα.IL-13Rα1.Fc) and pCAE802 (the DNA plasmid construct encoding IL-13Rα1.IL-4Rα.Fc) in DH10B cells were grown overnight in LB + ampicillin and the plasmid DNA was extracted using a Qiagen Endofree Mega Kit following the manufacturer's protocol. The concentration of the purified plasmid DNA was determined in a UV spectrophotometer and fluorometer. The plasmid DNA was also verified by digestion of aliquots with BbsI, XmnI and NcoI restriction enzymes. All restriction enzyme digest fragments corresponded to the predicted sizes in a 1% agarose gel.

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Forty 15 cm petri plates were seeded with CHO-K1/E1A cells at a density of 4×10^6 cells/plate. Plating media was Gibco Ham's F-12 w/10% Hyclone Fetal Bovine Serum (FBS) + penicillin/streptomycin and supplemented with glutamine. The following day each plate was transfected with 6 µg of pCAE801, or pCAE802, using Gibco Optimem and Gibco Lipofectamine in 12 ml volume, following the manufacturer's protocol. Four hours after adding the transfection mix to the cells 12 ml/plate of Optimem w/ 10% FBS was added. Plates were incubated at 37°C in a 5% CO₂ incubator overnight. The following day the media was removed from each plate and 25 ml expression media (Gibco CHO-S-SFM II w/ glutamine + 1mM sodium butyrate) was added. The plates were incubated at 37°C for 3 days.

After 3 days of incubation the media was removed from each plate and centrifuged at 400 rpm in a swinging bucket rotor to pellet cells. The supernatant was decanted into sterile 1L bottles and expressed protein was purified as described *infra*.

EXAMPLE 12: PURIFICATION OF IL-4Rα.IL-13Rα1.Fc AND IL-13Rα1.IL-4Rα.Fc PROTEIN FROM CULTURE MEDIA

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1. Purification of IL-4Rα.IL-13Rα1.Fc.

Human IL-4Rα.IL-13Rα1.Fc was transiently expressed in CHO cells and supernatants were harvested from plate transfections as described supra. Expression of the secreted protein was determined by a sandwich ELISA using goat anti-hIgG (y chain specific; Sigma 1-3382) and goat anti-hIgG (Fc specific)-FITC conjugate (Sigma F9512) capture and report antibodies, respectively. The yield ranged from 5.8 to 9.2 mg (average of 7.5 mg) per liter of conditioned media. CompleteTM protease inhibitor tablets (Roche 10 Diagnostics Corp.) were dissolved into the media (1 tablet/L). The conditioned media was sterile filtered (0.22 µm pore size) prior to loading onto a pre-equilibrated, 5 mL HiTrap® Protein A affinity column (Amersham Pharmacia Biotech) in Dulbecco's PBS buffer (Life Technologies), pH 7.4 at 4°C. The flow rate was ~1-2 mL/min. The column was extensively washed with PBS buffer to remove 15 nonspecifically bound proteins from the column. IL-4Rα.IL-13Rα1.Fc_was eluted using 20 mM sodium citrate, 150 mM NaCl, pH 3.5. The eluate was immediately neutralized by titrating with 1 M Tris-OH. The fractions containing protein were pooled and immediately dialyzed in PBS buffer, pH 7.4 at 4°C. The recovery from Protein A purification was 6.8 mg (73%). 20 IL-4Rα.IL-13Rα1.Fc was further purified by size exclusion chromatography using a superose 6 column (25 mL bed volume; Amersham Pharmacia Biotech) pre-equilibrated in PBS, 5% v/v glycerol, pH 7.4 at ambient temperature. The flow rate was 0.5 mL/min. Protein fractions were 25 assessed from a Coomassie stained non-reduced and reduced SDS-PAGE (Novex NuPAGE 4-12% Bis-Tris gels). Fractions were conservatively pooled to reduce the amount of aggregated protein. The overall yield was 51% (4.4 mg) with a purity of 97% as judged by SDS-PAGE. Purified IL-4Rα.IL-13Rα1.Fc was analyzed by non-reduced and reduced SDS-PAGE (4-12% Bis-Tris), analytical size exclusion chromatography (Tosohaas 30

TSKG4000SWXL), N-terminal sequencing, and immunoblotting with goat anti-hIgG-HRP conjugate (Promega W403B), and also mouse monoclonal anti-hIL-4R (R&D MAB230) followed by anti-mIgG-HRP conjugate (Promega W402B) as the secondary antibody.

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2. Purification of IL-13Rα1.IL-4Rα.Fc

Human IL-13Rα1.IL-4Rα.Fc was transiently expressed in CHO cells and supernatants were harvested from plate transfections as described supra. Expression of the secreted protein was determined by a sandwich ELISA using goat anti-hIgG (y chain specific; Sigma 1-3382) and goat anti-hIgG (Fc specific)-FITC conjugate (Sigma F9512) capture and report antibodies, respectively. The yield was 8.8 mg per liter of conditioned media. CompleteTM protease inhibitor tablets (Roche Diagnostics Corp.) were dissolved into the media (1 tablet/L). The conditioned media was sterile filtered (0.22 μm pore size) prior to loading onto a pre-equilibrated, 5 mL HiTrap® Protein A affinity column (Amersham Pharmacia Biotech) in Dulbecco's PBS buffer (Life Technologies), pH 7.4 at 4°C. The flow rate was ~1-2 mL/min. The column was extensively washed with PBS buffer to remove nonspecifically bound proteins from the column. IL-13Rα1.IL-4Rα.Fc was eluted using 20 mM sodium citrate, 150 mM NaCl, pH 3.5. The eluate was immediately neutralized by titrating with 1 M Tris-OH. The fractions containing protein were pooled and immediately dialyzed in PBS buffer, pH 7.4 at 4 °C. The recovery from Protein A purification was 3.8 mg (43%). IL-13Rα1.IL-4Rα.Fc was further purified by size exclusion chromatography using a superose 6 column (25 mL bed volume; Amersham Pharmacia Biotech) pre-equilibrated in PBS, 5% v/v glycerol, pH 7.4 at ambient temperature. The flow rate was 0.5 mL/min. Protein fractions were assessed from a Coomassie stained non-reduced and reduced SDS-PAGE (Novex NuPAGE 4-12% Bis-Tris gels). Fractions were

conservatively pooled to reduce the amount of aggregated protein. The overall yield was 17% (1.5 mg) with a purity of 95% as judged by SDS-PAGE. Purified IL-13Rα1.IL-4Rα.Fc was analyzed by non-reduced and reduced SDS-PAGE (4-12% Bis-Tris), analytical size exclusion chromatography (Tosohaas TSKG4000SWXL), N-terminal sequencing, and immunoblotting with goat anti-hIgG-HRP conjugate (Promega W403B), and also mouse monoclonal anti-hIL-4Rα (R&D MAB230) followed by anti-mIgG-HRP conjugate (Promega W402B) as the secondary antibody.

10 EXAMPLE 13: BLOCKING OF IL-4 AND IL-13 BY IL-4Rα.IL-13Rα1.Fc AND IL-13Rα1.IL-4Rα.Fc

Materials and Methods

TF1 Bioassay. TF1 cells were maintained in growth media (10ng/ml GM-CSF, RPMI 1640, 10% FBS, L-glutamine, Penicillin, Streptomycin). For the bioassay, cells were washed 2 times in assay media (as above but without GM-CSF) and then plated at 2 x 10⁵ cells in 50μl of assay media. The purified IL-4Rα.IL-13Rα1.Fc and IL-13Rα1.IL-4Rα.Fc proteins were diluted into assay media at a concentration of 40nM. 25ul of each of the traps was added to the cells. Either IL-13 or IL-4 were diluted to 40nM in assay media and then 2-fold dilution series in assay media were made. 25μl of either IL-13 or IL-4 was then added to the wells containing the cells and the traps. Cells were then incubated at 37°C, 5% CO₂ for ~70 hrs. The extent of TF1 cell proliferation was measured by the MTS assay according to the manufacturer's protocol (Promega, Inc.).

RESULTS

The ability of the IL-4Rα.IL-13Rα1.Fc and IL-13Rα1.IL-4Rα.Fc traps to block both human IL-13 and human IL-4 activity was measured in the TF1

bioassay described *supra*. IL-13 stimulates proliferation of TF1 cells, with half-maximal growth at a concentration of 0.2nM. Addition of either IL-4Rα.IL-13Rα1.Fc or IL-13Rα1.IL-4Rα.Fc trap at a concentration of 10nM blocks IL-13-induced growth up to ~2nM (Figure 33). At an IL-13 concentration of ~4-5 nM the growth of TF1 cells is inhibited by 50%. TF1 cells are more sensitive to IL-4, which stimulates their proliferation with half-maximal growth at ~0.02nM. Addition of either IL-4Rα.IL-13Rα1.Fc or IL-13Rα1.IL-4Rα.Fc at a concentration of 10nM blocks IL-4-induced growth up to ~1nM (Figure 34). At an IL-4 concentration of ~3-4 nM the growth of TF1 cells is inhibited by 50%. These results show that both IL-4Rα.IL-13Rα1.Fc and IL-13Rα1.IL-4Rα.Fc can block the ability of both IL-13 and IL-4 to stimulate cellular responses.

EXAMPLE 14: BLOCKING OF INJECTED IL-1 BY IL-1 TRAP IN VIVO

IL-1 is a pro-inflammatory cytokine. Systemic administration of IL-1 has been shown to elicit acute responses in animals, including transient hyperglycemia, hypoinsulinemia, fever, anorexia, and increased serum levels of interleukin-6 (IL-6) (Reimers, 1998). Since mice are responsive to both murine and human IL-1, human IL-1 can be used and *in vivo* binding effects of human specific IL-1 antagonists can be evaluated. This acute mouse model was used to determine the ability of a human IL-1 trap to antagonize the *in vivo* effects of exogenously administered human IL-1. This provides a rapid indication of *in vivo* efficacy of the human IL-1 trap and can be used as an assay to help molecule selection.

Experimental Design:

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Mice were given subcutaneous injections of human IL-1 (0.3 µg/kg).

Twenty-four hours prior to human IL-1 injection, the animals were pretreated with either vehicle or 150-fold molar excess of human IL-1 trap (0.54 mg/kg). Two hours prior to sacrifice (26 hrs), the mice were given a

second injection of human IL-1 (0.3 µg/kg). Blood samples were collected at various time points and sera were assayed for IL-6 levels.

RESULTS

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Exogenous administration of human IL-1 resulted a dramatic induction of serum IL-6 levels. At 150-fold molar excess, the human IL-1 trap completely blocked the IL-6 increase (Figure 35). Furthermore, the effects of the human IL-1 trap persisted for at least another 24 hours, preventing an IL-6 increase even when IL-1 was re-administered (Figure 35). Such long-lasting efficacy suggests that daily injection of an IL-1 trap may not be necessary for chronic applications.

EXAMPLE 15: EVALUATING THE ABILITY OF AN IL-4 TRAP TO BLOCK THE PHYSIOLOGICAL RESPONSES TO HUMAN IL-4 IN CYNOMOLOGUS MONKEYS.

Systemic administration of human IL-4 elicits systemic responses in Cynomologus monkeys (Gundel et al., 1996). Thus, the effectiveness of the IL-4 trap in blocking human IL-4 can be demonstrated by measuring these responses.

Experimental Design:

The experiment consisted of 3 parts: human IL-4 + vehicle (part 1), human IL-4 + IL-4 Trap (part 2), and human IL-4 + vehicle (part 3).

Human IL-4 (25 μg/kg) was injected subcutaneously twice daily for 4 days and IL-4 Trap (8 mg/kg) and vehicle were given intravenously daily for 5 days, beginning 1 day prior to human IL-4 administration. Whole blood was collected daily for flow cytometry analysis for CD16 and plasma was obtained to assay for the cytokine monocyte chemotactic protein 1 (MCP-1).

CD16 and MCP-1 are markers of IL-4-mediated inflammation in both humans and monkeys.

RESULTS

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In the presence of human IL-4, MCP-1 increased 2.5-fold and was significantly blocked by the IL-4 Trap (Figure 36A). Similarly, the decrease in the percent of CD16 positive lymphocytes in peripheral blood was attenuated by the IL-4 trap (Figure 36B). After a rest period, the monkeys were re-injected with human IL-4 and the responsiveness of the animals to human IL-4 was re-confirmed (Figures 36A and 36B), suggesting that inhibition of the MCP-1 and CD 16 responses is specifically mediated by the IL-4 trap.

15 EXAMPLE 16: THE EFFECTS OF IL-4 TRAP ON 1L-4-INDUCED IgE SECRETION.

It has been shown that injection of anti-mouse IgD antibody stimulates an IL-4-mediated IgE increase in normal mice. This model has been widely used to evaluate IL-4 antagonists, such as soluble IL-4 receptor and anti-IL-4 monoclonal antibodies (Sato et al., 1993). We decided to use this model to evaluate the ability if the IL-4 trap to block IL-4-mediated increases of IgE.

25 Experimental design:

BALB/C mice injected with anti-mouse IgD (100µl/mouse, s.c.) were randomly divided into 3 groups. Each received (on days 3-5) either vehicle, murine IL-4 trap (1 mg/kg, s.c.), or a monoclonal antibody to mouse IL-4 (1 mg/kg, s.c.). Serum was collected at various time points and assayed for IgE levels.

RESULTS

Treatment with the murine IL-4 trap or the mouse IL-4 antibody both significantly antagonized the IL-4-mediated IgE increase in this mouse model (Figure 37). This suggests that the murine IL-4 trap binds murine IL-4 and antagonizes physiological responses elicited by endogenous IL-4 *in vivo*.

The present invention is not to be limited in scope by the specific

10 embodiments described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description and accompanying figures. Such modifications are intended to fall within the scope of the appended claims.

WE CLAIM:

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- 1. An isolated nucleic acid molecule encoding a fusion polypeptide capable of binding a cytokine to form a nonfunctional complex comprising:
- a) a nucleotide sequence encoding a first fusion polypeptide component comprising the amino acid sequence of the cytokine binding portion of the extracellular domain of the specificity determining component of the cytokine's receptor;
- b) a nucleotide sequence encoding a second fusion polypeptide component comprising the amino acid sequence of the cytokine binding portion of the extracellular domain of the signal transducing component of the cytokine's receptor; and
- c) a nucleotide sequence encoding a third fusion polypeptide component comprising the amino acid sequence of a multimerizing component.
 - 2. The nucleic acid molecule of claim 1, wherein the nucleotide sequence encoding the first component is upstream of the nucleotide sequence encoding the second component.
 - 3. The nucleic acid molecule of claim 1, wherein the nucleotide sequence encoding the first component is downstream of the nucleotide sequence encoding the second component.

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4. The isolated nucleic acid molecule of claim 1, wherein the cytokine receptor is the receptor for a member of the hematopoietin family of cytokines selected from the group consisting of interleukin-2, interleukin-3, interleukin-4, interleukin-5, interleukin-6, interleukin-7, interleukin-9, interleukin-11, interleukin-13, interleukin-15, granulocyte macrophage colony stimulating factor, oncostatin M, and leukemia inhibitory factor and cardiotrophin-1

5. The isolated nucleic acid molecule of claim 1, wherein the cytokine receptor is the receptor for a member of the interferon family of cytokines selected from the group consisting of IFN-gamma, IFN-alpha, and IFN-beta.

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6. The isolated nucleic acid molecule of claim 1, wherein the cytokine receptor is the receptor for a member of the immunoglobulin superfamily of cytokines selected from the group consisting of B7.1 (CD80) and B7.2 (B70).

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7. The isolated nucleic acid molecule of claim 1, wherein the cytokine receptor is the receptor for a member of the TNF family of cytokines selected from the group consisting of TNF-alpha, TNF-beta, LT-beta, CD40 ligand, Fas ligand, CD 27 ligand, CD 30 ligand, and 4-1BBL.

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- 8. The isolated nucleic acid molecule of claim 1, wherein the cytokine receptor is the receptor for a member of the TGF-β/BMP family selected from the group consisting of TGF-β1, TGF-β2, TGF-β3, BMP-2, BMP-3a, BMP-3b, BMP-4, BMP-5, BMP-6, BMP-7, BMP-8a, BMP-8b, BMP-9, BMP-10, BMP-11, BMP-15, BMP-16, endometrial bleeding associated factor (EBAF), growth differentiation factor-1 (GDF-1), GDF-2, GDF-3, GDF-5, GDF-6, GDF-7, GDF-8, GDF-9, GDF-12, GDF-14, mullerian inhibiting substance (MIS), activin-1, activin-2, activin-3, activin-4, and activin-5.
- 9. The isolated nucleic acid molecule of claim 1, wherein the cytokine receptor is the receptor for a cytokine selected from the group consisting of interleukin-1, interleukin-10, interleukin-12, interleukin-14, interleukin-18 and MIF.
- 30 10. The isolated nucleic acid molecule of claim 1, wherein the multimerizing component comprises an immunoglobulin derived domain.

11. The isolated nucleic acid molecule of claim 10, wherein the immunoglobulin derived domain is selected from the group consisting of the Fc domain of IgG, the heavy chain of IgG, and the light chain of IgG.

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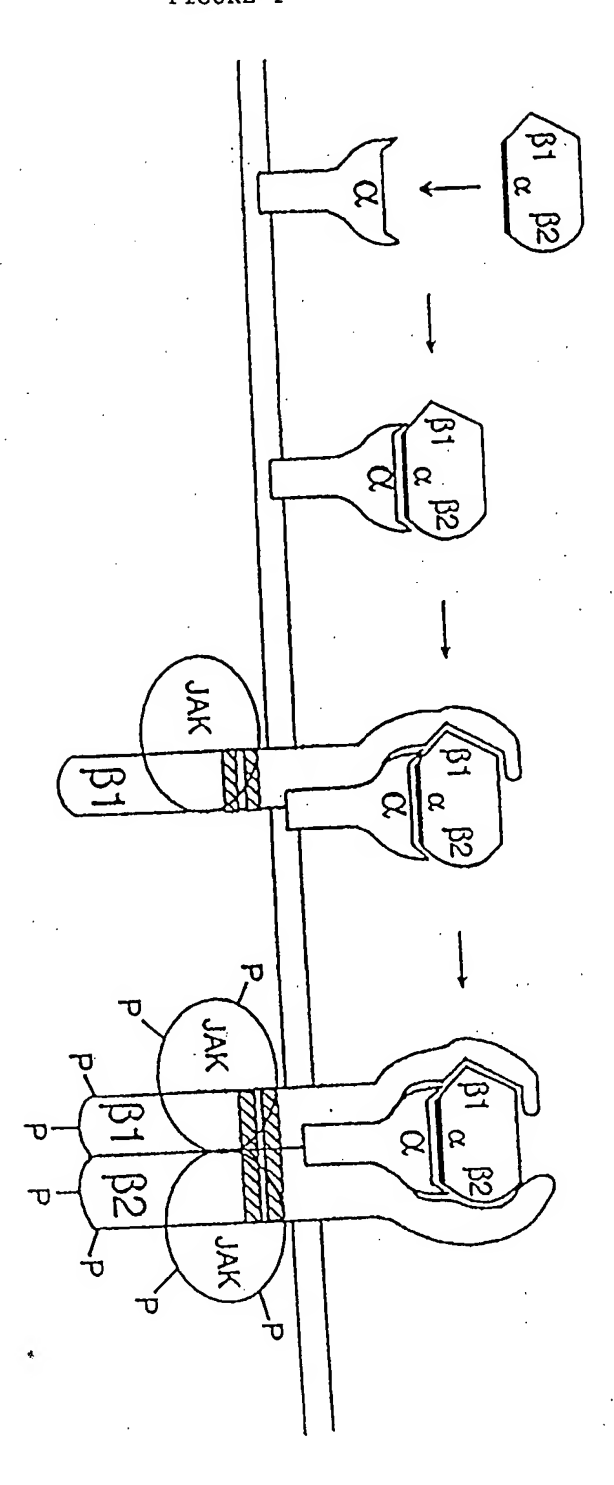
- 12. A fusion polypeptide encoded by the isolated nucleic acid molecule of claim 1.
- 13. A composition capable of binding a cytokine to form a nonfunctional complex comprising a multimer of the fusion polypeptide of claim 12.
 - 14. The composition of claim 13, wherein the multimer is a dimer.
- 15 15. A vector which comprises the nucleic acid molecule of claim 1.
 - 16. An expression vector comprising a nucleic acid molecule of claim 1, wherein the nucleic acid molecule is operatively linked to an expression control sequence.

- 17. A host-vector system for the production of a fusion polypeptide which comprises the expression vector of claim 16, in a suitable host cell.
- 18. The host-vector system of claim 17, wherein the suitable host cell is a bacterial cell, yeast cell, insect cell, or mammalian cell.
 - 19. The host-vector system of claim 17, wherein the suitable host cell is <u>E. coli</u>.
- 30 20. The host-vector system of claim 17, wherein the suitable host cell is a COS cell.

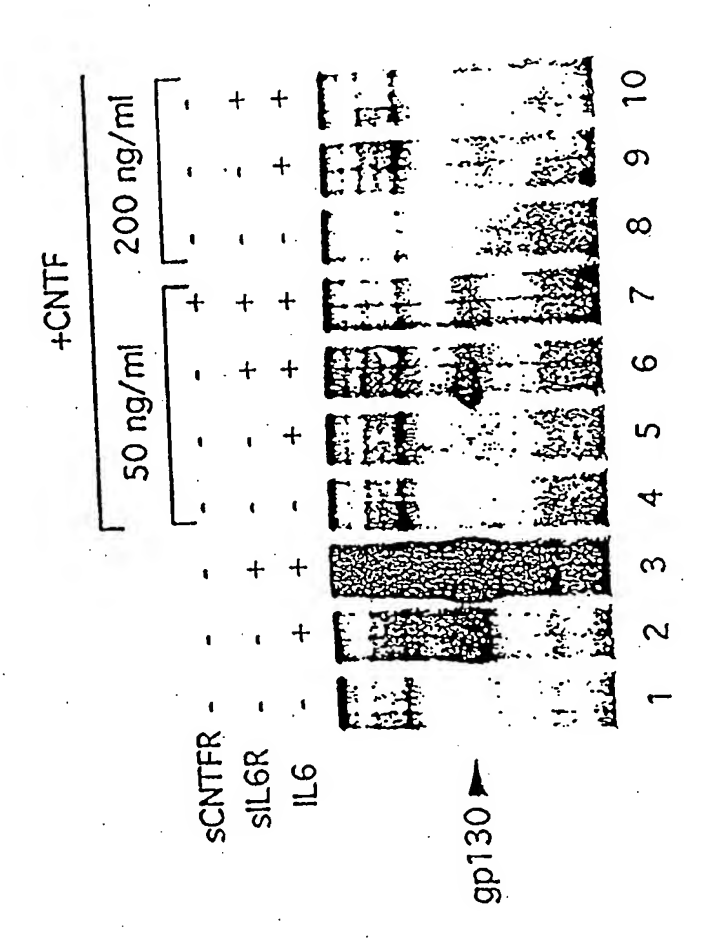
21. The host-vector system of claim 17, wherein the suitable host cell is a CHO cell.

- 22. The host-vector system of claim 17, wherein the suitable host cell is a 293 cell.
 - 23. The host-vector system of claim 17, wherein the suitable host cell is a BHK cell.
- 10 24. The host-vector system of claim 17, wherein the suitable host cell is a NS0 cell.
- 25. A method of producing a fusion polypeptide which comprises growing cells of the host-vector system of claim 17, under conditions
 15 permitting production of the fusion polypeptide and recovering the fusion polypeptide so produced.

1/74 FIGURE 1



2/ 74 FIGURE 2



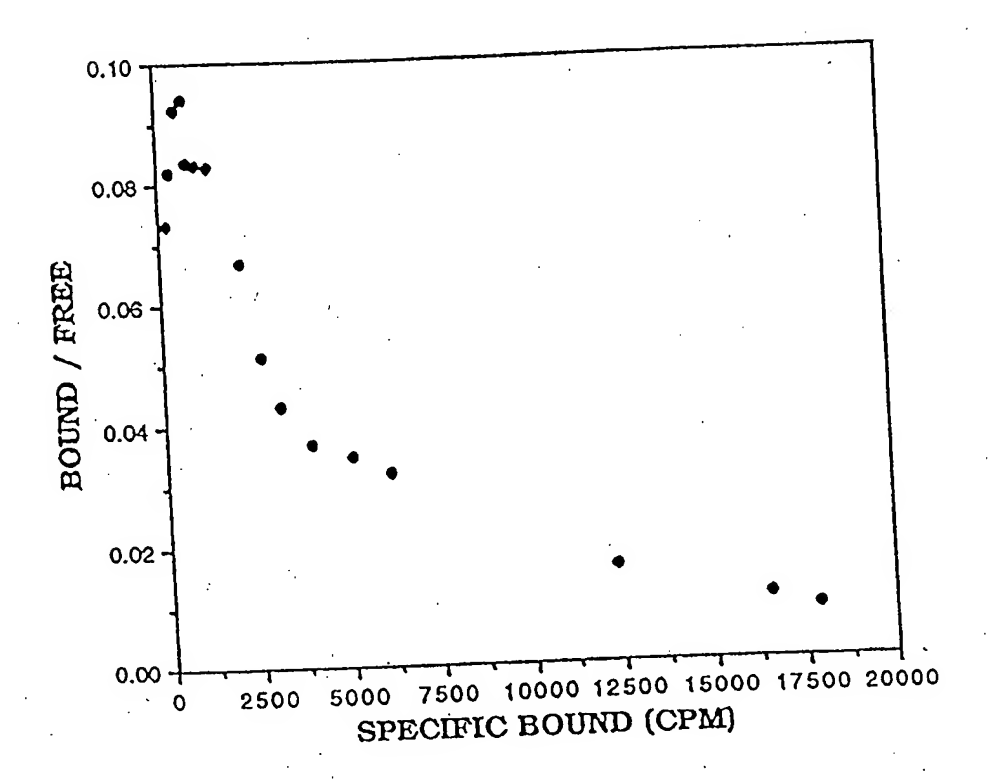


Figure 4

Amino acid sequence of human gp130-Fc-His6

Sequence Range	e: 1 to 861				
10	20	30	40	50 *	60 *
* MVTLQTWVVQALF	* TELTTES TGELL	.DPCGYISPESI	PVVQL HSNF	TAVCVLKEKCMI	YFHV
WAIPÕIMAAÕYDE			100	110	120
70 *	* 80	90 *	*	* .	* ************************************
* NANYIVWKTNHFT	IPKEQYT IINR	rassvtftdia	SLNIQ LTCI	41P.L.E.GÖTFÖNA	IGIII
130	140	150	160	170	180 *
	* SCIVNEGK KMRC	* EWDGGRETHLE	TNFTL KSE	WATHKFADCKAK	RDTPT
ISGLPPEKPKNL	SCIAMERY WING			230	240
190	200	210	220	*	*
SCTVDYSTVYFV	NIEVWVEA ENAI	GKVTSDHINF	DBAAKA KBR	IPPHNLSVINSEE	PSSID
250	260	270	280	290	300 *
	* VIILKYNIQ YRT	* KDÁSTWSOIPP	EDTAST RS	SFTVQDLKPFTE	YVFRIR
KLTWTNPSIKSV	TIPKINIÖ IVI		340	350	360
310	· 320 *	330 *	*	*	*
CMKEDGKGYWSI	DWSEEASGI TYE	DRPSKAPSFWY	rkidpsh tQ	GYRTVQLVWKTL	PPFEAM
370	380	390	400	410 *	420
•	* RWKSHLQNY TVI	* JATKLTVNLTN	DRYLATL TV	/RNLVGKSDAAVI	TIPACD
GKILDAEALDI			460	470	480
430	440 *	450 *	*	*	*
FQATHPVMDLE	(AFPKDNMLW VE	WTTPRESVKKY	ILEWCVL S	DKAPCITDWQQE	DGTVRKI
4.90	500	510	520	530 *	540 *
	* CYLITVTPVY AD	* KOPGSPESIKA	YLKQAPPS K	GPTVRTKKVGK	IEAVLEWD
YLRGNLAESK			580	590	600
550 *	*	570 *	*	*	* * * * * * * * * * * * * * * * * * * *
QLPVDVQNGF	'IRNYTIFYRT I	IGNETAVNVDS	SHTEYTLS S	SLTSDTLYMVRM	
610	620	630	640	650 *	· 660
•	rpkfaQgeies g	t * EPKSCDKTHTC	T PPCPAPEL	LGGPSVFLFPPK	PKDTLMIS
KDGPEFTFT.	LAKLYÖGETED O		700	710	720
67		690 *	*	*	*
RTPEVTCVV	*	NWYVDGVEVH	NAKTKPREE	OYNSTYRVVSVI	<u>`.T.A PHODAI</u>
73	0 740	750	760	770	780
1.3	, - -	*	*	*	

HYTOKSLSLSPGKHHHHHH.

FIGURE 4 continued

NGKEYKCKVSNKALPAPIEK TISKAKGOPREPOVYTLPPS RDELTKNOVSLTCLVKGFYP

790 800 810 820 830 840

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6/74 FIGURE 5

The amino acid sequence of human IL-6Rα-Fc

	•				•
Sequence Range:	1 to 594				
10	20	30	40	50 *	60 *
* MVAVGCALLAAĻLA	* A <i>PGAA</i> L APRRO	* CPAQEVARGVL	TSLPG DSVTI	TCPGVEPEDN	WHVTA
MANAGCADITATION				110	120
70	80	90 *	100	*	*
* VLRKPAAGSHPSRW	AGMGRR LLLR	SVQLHDSGNYS	CYRAG RPAG	TVHLLVDVPPI	EEPQLS
130	140	150	160	170	180
	•	de	*	COVCORSOKE *	SCOLAV
* CFRKSPLSNVVCEW	GPRSTP SLTT	KAVLLVRKFQ	NSPAED FQEP	CÖTPÖRPÖIG	DCQ
190	200	210	220	230	240
		*	* NTM	massarnprwi	SVTWOD
PEGDSSFYIVSMC	vassvgs kfsi	KTQTFQGCGIL	WADDAY MII	TVALTORI	
250	260	270	280	290	300
		*	* Will Daw	sgr.RHVVOLRA	OEEFGQ
* PHSWNSSFYRLRF	ELRYRAE RSK	TETTWMVKDLY	SHHCAIR DYII		
310	320	330	340	350	360 *
	*	*	TATINGTOD DNI	LFRDSANATS	LPVQDAG
.* GEWSEWSPEAMGT	PWTESRS PPA	ENEVSTPMQA	Billion 200		
370	380	390	400	410	420 *
*† EPKSCDKTHTCP	† *	* homer EDDKDK	TYPLMISR TPI	EVICVVVDVSI	EDPEVKE
EPKSCDKTHTCP	PCPAPELL GG	BALDLLIKEY			
430	440	450	460	470	480 *
* NWYVDGVEVHNA	*	* ************************************	T.HODWLN GK	EYKCKVSNKA	LPAPIEKT
NWYVDGVEVHNA	KTKPREEO IN	STIRVADADA			
490	500	510	520	530	540 *
* ISKAKGOPREPO	*	* OP.TSMOMSE	LVKGFYPS DI	AVEWESNGOR	ENNYKTTP
<u> ISKAKGOPREP</u>	OVYTLPPSR DI	SPIKNOASDIC	DVXCFFFF		
550	560	570	580	590 *	
* PVLDSDGSFFL	*	*	* линат.нин Ү	TOKSLSLSPGI	<u> </u>
PVLDSDGSFFL	YSKLTVDKS R	MOOPHALDEDI	1 P Y Y Y 4-15 P 4-25 - 24 1 5 2		



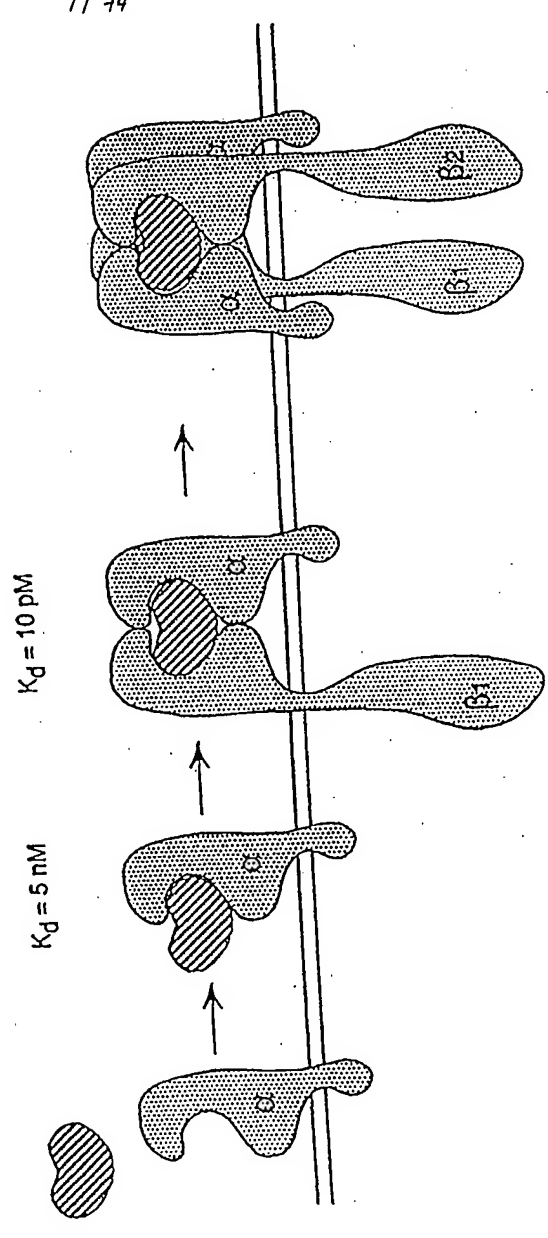
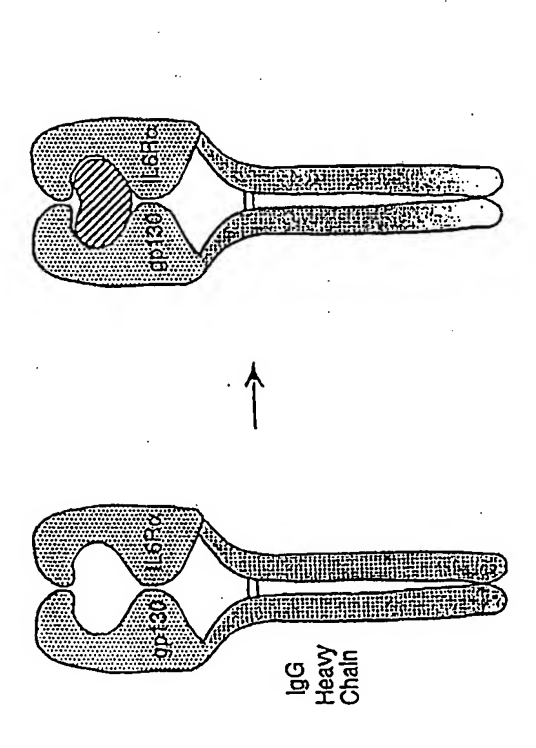
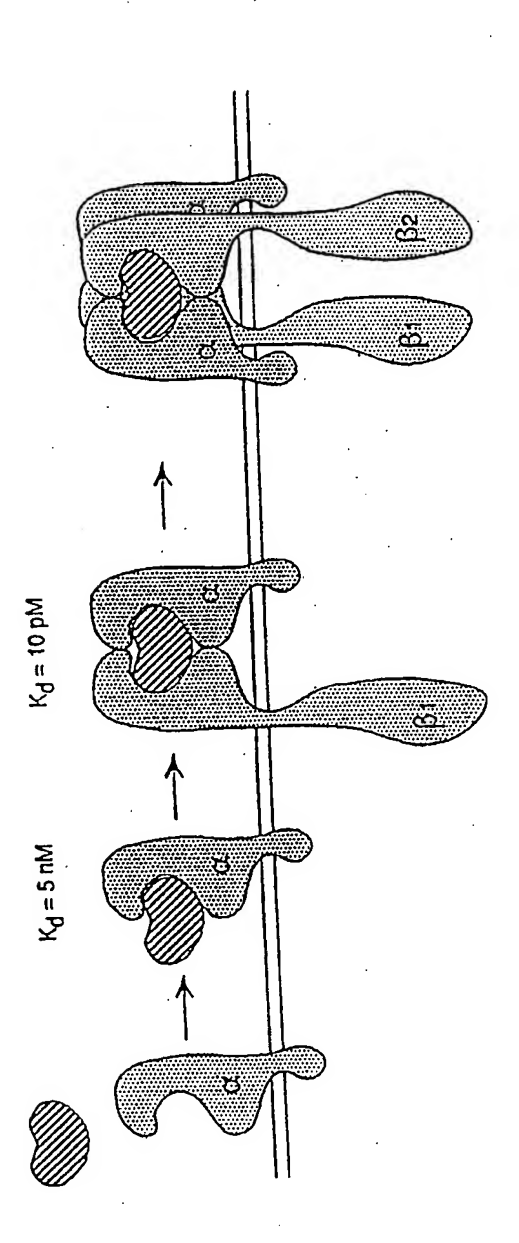
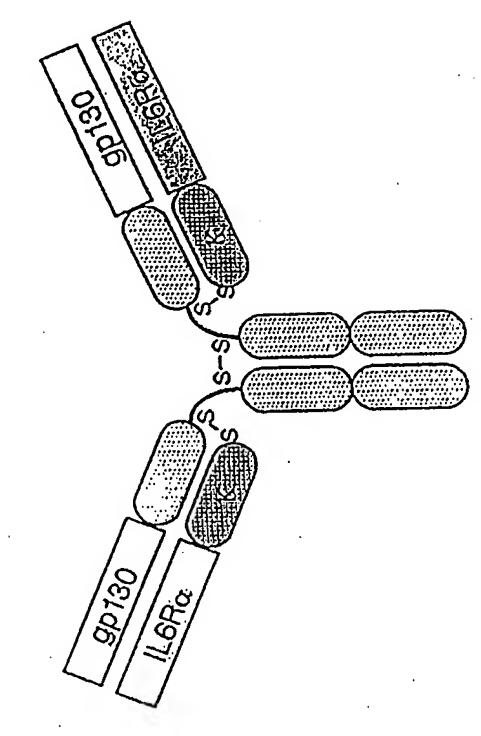


FIGURE 7
Heterodimeric Receptor-Based Ligand Trap





Immunoglobulin Heavy/Light Chain Receptor Fusions



10/74 FIGURE 9

Amino acid sequence of gp130-Cy1

Sequence R	Range: 1	to 952		٠		
10		20	30	40	. 50 *	60 *
VVWTO,ITVM	* DALFIFLT	* YES TGELLDI	* PCGYISPES	PVVQL HSNI	FTAVCVLKEKCMD	YFHV
7(80	90	100	110	120
NANYIVWKT	* NHFTIPKE	YT IINRTA	SSVTFTDIA	SLNIQ LTC	NILTFGQLEQNVY	GITI
13		140	150	160	. 170	180
ISGLPPEKP	* KNLSCIVN	EGK KMRCEW	DGGRETHLE	etnftl kse	WATHKFADCKAKI	UTPT
. 19	0	200	210	220	230	240
SCTVDYSTV	* /YFVNIEVY	IVEA ENALGI	CVTSDHINE	DPVYKV KPI	NPPHNLSVINSEE:	LSSIL
25		260	270	280	290	300
KLTWTNPS:	* IKSVIILK	* (NIQ YRTKD)	ASTWSQIPP	EDTAST RS	SFTVQDLKPFTEY	VFRIR
	10	320	330	340	350	360 *
CMKEDGKG	* YWSDWSEE	* ASGI TYEDR	* PSKAPSFWI	KIDPSH TQ	GYRTVQLVWKTLI	PFEAN
3	70.	380	390	400 *	410	420 *
GKILDYEV	* TLTRWKSH	LQNY TVNAI	KLTVNLTN	DRYLATL 'T	/RNLVGKSDAAVL	ripacd
	130	440	450 *	460 *	470 *	480 *
FQATHPV	* MDLKAFPKI	NMLW VEWT	rpresvkky	ILEWCVL S	DKAPCITDWQQED	GTVHRT
	490	500	510	520 *	530 *	540 *
YLRGNLA	* ESKCYLIT	* VTPVY ADGP	GSPESIKAY	KLKQAPPS K	GPTVRTKKVGKNI	EAVLEWD.
	550	560	570	580	590 *	600 *
QLPVDVQ	* NGFIRNYT	* IFYRT IIGN	ETAVNVDS:	SHTEYTLS S	SLTSDTLYMVRMA	AYTDEGG
	610	620	630	640	650 *	660
KDGPEFT	* rfttpkfa(XGEIES G <u>AS</u>	rkgpsvfpl	APSSKSTS (GGTAALGCLVKDY	FPEPVTV
	670	680	690	700 *	710	720
SWNSGA	* LTSGVHTF	PAVLOS SGL	YSLSSVVTV	PSSSLGTO	TYICNVNHKPSN	rkydkkyb
	730	740	750	760	770	780
<u>PKSCDK</u>	* THTCPPCP	* APELLG GPS	* SVFLFPPKPI	KDTLMISRT	PEVTCVVVDVSH	EDPEVKFI

11/74
FIGURE 9 continued

790 8t 810 820 8: 840

WYVDGVEVHNAKTKPREEOY NSTYRVVSVLTVLHODWLNG KEYKCKVSNKALPAPIEKTI

850 860 870 880 890 900

SKAKGOPREPOVYTLPPSRD ELTKNOVSLTCLVKGFYPSD IAVEWESNGOPENNYKTTPP

910 920 930 940 950

12/ 44 FIGURE 10

Amino acid sequence or gp130∆3fibro

Sequence Range: 1 to 332 MVTLQTWVVQALFIFLTTES TGELLDPCGYISPESPVVQL HSNFTAVCVLKEKCMDYFHV NANYIVWKTNHFTIPKEQYT IINRTASSVTFTDIASLNIQ LTCNILTFGQLEQNVYGITI .160 ISGLPPEKPKNLSCIVNEGK KMRCEWDGGRETHLETNFTL KSEWATHKFADCKAKRDTPT SCTVDYSTVYFVNIEVWVEA ENALGKVTSDHINFDPVYKV KPNPPHNLSVINSEELSSIL KLTWTNPSIKSVIILKYNIQ YRTKDASTWSQIPPEDTAST RSSFTVQDLKPFTEYVFRIR CMKEDGKGYWSDWSEEASGI TYEDRPSKAPSG

•

FIGURE 11

Amino acid sequence of J-CH1

FIGURE 12

Amino acid sequence of Cy4

Sequence	Range: 1	to 330				
:	1.0	20	30	40 *	50 *	60 *
SGASTKGP	SVFPLAPCSI	RST SESTAAL	GCLVKDYFP	EPVT VSWNSG	ALTSGVHTFP	PAVLQ
	70	80	90	100	110	120
SSGLYSLS	* SVVTVPSSS	LGT KTYTCN	MHKPSNTKV	DKRV ESKYGE	PCPSCPAPE	PLGGP
· 1	.30	140	150	160	170	180
SVFLFPPK	KPKDTLMISR	TPE VTCVVV	DVSQEDPEVQ	FNWY VDGVE	VHNAKTKPRE!	EQFNS
	L90	200	210	220	230	240 *
TYRVVSVI	LTVLHQDWL <i>N</i>	GKE YKCKVS	nkglpssiei	KTISK AKGQP	REPQVYTLPP	SQEEM
	250	260	270	280	290	300 *
TKNQVSL	TCLVKGFYP	SDIA VEWESN	IGQPENNYK T	TPPVL DSDGS	FFLYSRLTVI)KSRWQ
	310	320	330 *			
EGNVFSC	SVMHEALHN	HYTQ KSLSL	SLGK*			

FIGURE 13

Amino acid sequence of k-domain

Sequence Range: 1 to 108

70 80 90 100

DSKDSTYSLSSTLTLSKADY EKHKVYACEVTHQGLSSPVT KSFNRGEC*

FIGURE 14

Amino acid sequence of λ -domain:

Sequence Range: 1 to 107

10 20 30 40 50 60 * *

SGPKAAPSVTLFPPSSEELQ ANKATLVCLISDFYPGAVTV AWKADSSPVKAGVETTPSK

70 80 90 100 * *

QSNNKYAASSYLSLTPEQWK SHRSYSCQVTHEGSTVEKTV APTECS*

17/ 74 FIGURE 15

Amino acid sequence of the soluble IL-bkx domain

Sequence Range:	1 to 360				
10	20	30	40	50 *	60 *
MVAVGCALLAALLA	APGAAL APR	RCPAQEVARGV	LTSLPG DSVI	LTCPGVEPEDI	WHVTAN
7:0	80	90	100	110	120
* VLRKPAAGSHPSRW	agmgrr lli	RSVQLHDSGNY	SCYRAG RPA	STVHLLVDVPP	EEPQLS
130	140	150	160 *	170	180
* CFRKSPLSNVVCE	GPRSTP SL	rtkavllvrkf(ONSPAED FQE	PCQYSQESQKE	SCQLAV
190	200	210	220 *	230	240
PEGDSSFYIVSMC	vassvgs kf	SKTQTFQGCGI	LQPDPPA NIT	'VTAVARNPRW	LSVTWQD
250	260	270	280 *	290 *	300
* PHSWNSSFYRLRF	ELRYRAE RS	KTFTTWMVKDI	QHHCVIH DAV	NSGLRHVVQLR	AQEEFGQ
310	320	330	340	350 *	360 *
* GEWSEWSPEAMGT	*PWTESRS P	PAENEVSTPMQ	ALTTNKDD DN	ILFRDSANATS	SLPVQDAG

FIGURE 16

Amino acid sequence of the soluble IL-6kx313 domain

Sequence	Range: 1	to 3	15		•	
	10	20.	30	40	50 *	60 *
MVAVGCAL	LAALLAAPG	AAL A	APRRCPAQEVARGV	LTSLPG T	SVTLTCPGVEPE	WHVTANC
	70	80	90	100	110	120 *
VLRKPAAG	* SHPSRWAGM	GRR	LLLRSVQLHDSGNY	SCYRAG I	RPAGTVHLLVDVP	PEEPQLS
	.30	140	150 *	160 *	. 170 *	180
CFRKSPLS	NVVCEWGPF	RSTP	SLTTKAVLLVRKF(2NSPAED	FQEPCQYSQESQK	FSCQLAV
	190	200	210	220	230	240
PEGDSSF	YIVSMCVAS	SVGS	KFSKTQTFQGCGI	LQPDPPA	NITVTAVARNPRV	ILSVTWQD
	250	260	270	280	290	300 *
PHSWNSS	FYRLRFELR	YRAE	RSKTFTTWMVKDL	OHHCVIH	DAWSGLRHVVQLI	RAQEEFGQ
	310					
			•			

FIGURE 17

$$(gpx-C\gamma 1)^2 - \frac{1}{200} \frac{(gpx-C\gamma 1)^2 \cdot (6R\kappa)^2}{(gpx-C\gamma 1)^2 \cdot (6R\kappa)^2}$$

100

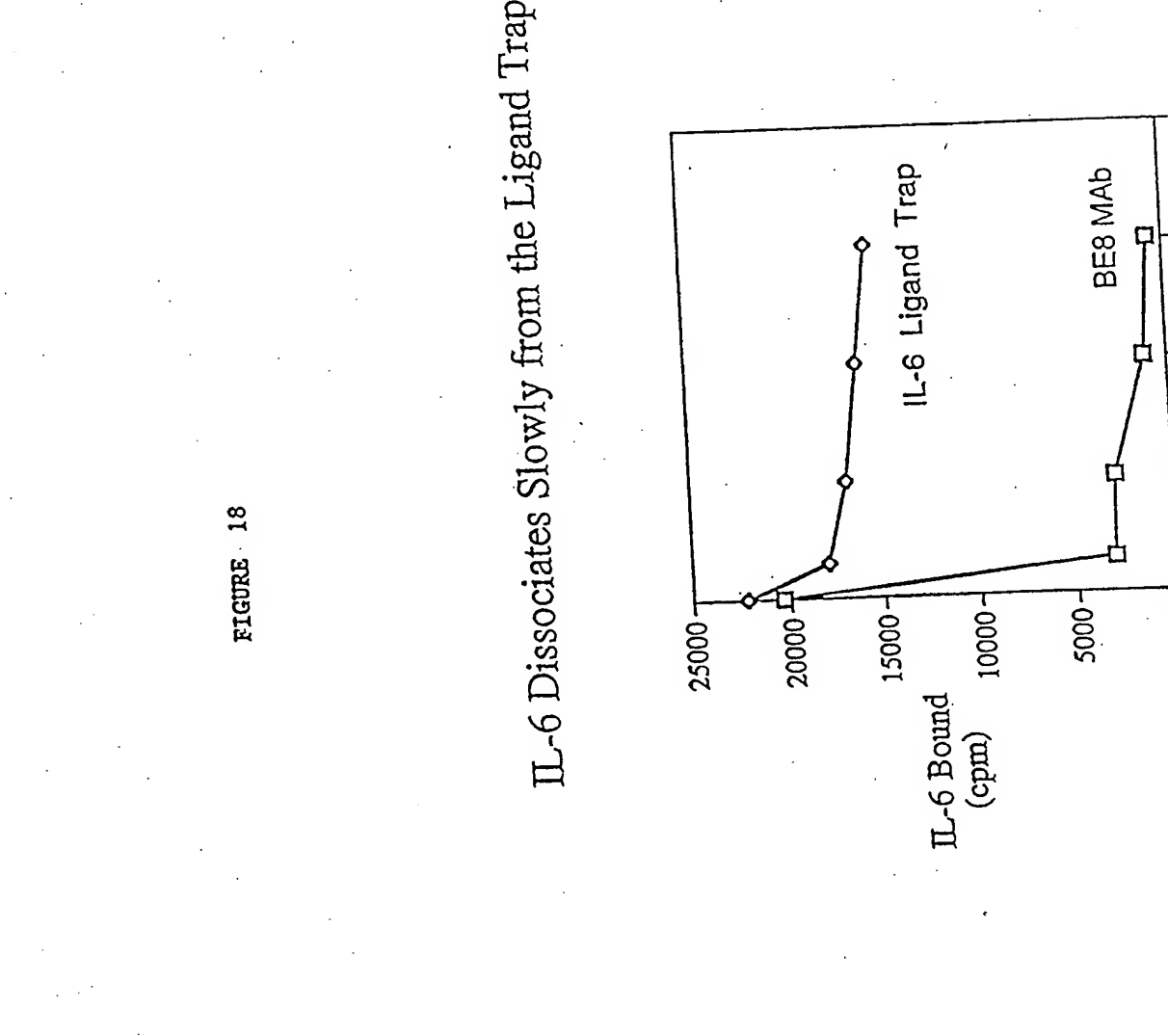
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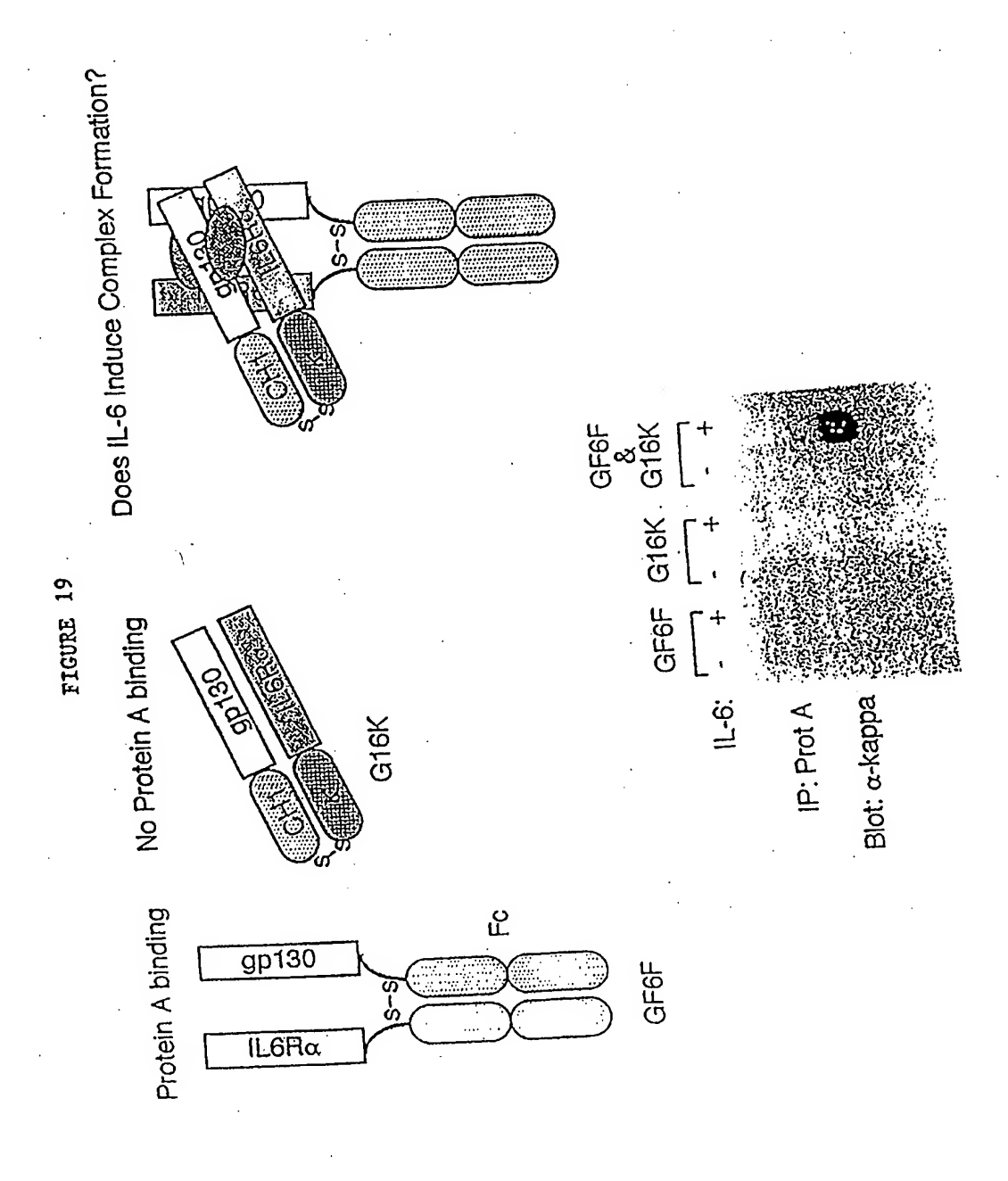
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Days

FIGURE







gp130Δ3fibro-Cγ1•IL-6Rα-j-κ gp130-J-CH1•IL-6Rα-j-κ gp130-Cγ4•IL-6Rα-κ gp130-Cγ1•IL-6Rα-κ gp130-Fc•IL-6Rα-Fc mAb B-E8 1000 100_ 20 cell proliferation assay FIGURE. of ligand trap over D -01 029-072A

		1	0			20		•	30		*	4	. O	*		
ATG Met	* GTG Val	AAG Lys	* CCA Pro	TCA Ser	TTA Leu	CCA Pro	TTC Phe	ACA Thr	TCC Ser	CTC Leu	TTA Leu	TTC Phe	CTG Leu	CAG Gln	CTG Leu>	
50			60		<u>.</u>	T	70	*		80		*	90		*	
* CCC Pro	CTG Leu	* CTG Leu	GGA Gly	GTG Val	GGG	CTG Leu	AAC Asn	ACG Thr	ACA Thr	ATT	CTG Leu	ACG Thr	CCC Pro	AAT Asn	GGG Gly>	•
. 1	00			110			120)	•	1	30	*		140		
AAT Asn	* GAA Glu	* GAC Asp	ACC Thr	* ACA Thr	GCT Ala	GAT Asp	TTC Phe	TTC Phe	CTG	ACC Thr	ACT Thr	ATG Met	CCC Pro	ACT	GAC Asp	>
	150		•	1	.60 .	1	k	170		*	180) *	*	1	L90 *	
TC(CTC	AGI Ser	GTT Val	TCC LSex	ACT Thi	CTC Lev	ı Pro	C CTO	ı Pro	A GAC	i Val	r CAC	n Cys	s Phe	r GTG e Val	>
		200			210) ·	4		220		*	230		*	240) •
TT Ph	* C AA' e As:	r GT(n Va	C GAG	TAC U TY	C ATO	G AA t As	т TG n Су	C AC	T TG r Tr	G AA p As	C AG n Se	C AG	C TC r Se	T GA r Gl	G CCC	C >>
			250			260		*	27	*	*		280		*	•
CA G3		T AC	* C AA ir As	C 000	 	יוני כיונ	יב כז	AT TA is Ty	T TO	G TA	AC AA	AG AA YS AS	C TC	G GA	AA TA sp As	T n>
290		· .		00	,	r	310		.	320	*	*		30 *	*	
GZ A	AT AI sp Ly	AA GT ys Va	rc ca al G		. a m	30 A	~C C	AC T	AT C	TA T' eu P	TC To	CT G er G	AA G	AA A' lu I	TC AC	r>
	340			35	0	*	3	60		*	370 *		*	38	0 *	
T	* CT G er G	GC T	* GT C Ys G		ma a	R R A	א תת	AC C	AG A	TC C	AC C	TC T Leu T	YAC C	AA A Sln T	CA T	TT he>
	3	90	•		400		*	41	LO *	1		120 *		*	430	
	GTT C Val V	* GTT C Val C	CAG C	* CTC C Leu C	AG C Sln <i>R</i>	120 (ירא (ccc (AAS	acc 2	AĠG Z	AGA (Arg (CAG (Gln)	GCC A Ala '	ACA C	AG Sln>
		44	40			150			46	0	*	. 4	70 *		* .	180 *
	* ATG ' Met	CTA /	* AAA (Lys	omo (* CAG . Gln .	AAT Asn	CTG Leu	CTC	ATC. Ile	Pro Pro	TGG Trp	GCT Ala	CCA Pro	GAG Glu	AAC (Asn	CTA Leu>
			49	0	•	5	00		*	510		*	5 2	.O *	· *	
	ACA Thr	* CTT Leu	CAC His	* AAA Lys	CTG Leu	AGT Ser	GAA Glu	TCC Ser	CAG Gln	CTA Leu	GAA Glu	CTG Leu	AAC Asn	TGG Trp	AAC Asn	AAC Asn>
ŗ	530		*	540		*		50 *	*		560 ·		*	570		*
	* AGA Arg	TTC Phe	TTG Leu	AAC Asn	CAC His	TGT Cys	TTG Leu	GAG Glu	CAC His	TTG Leu	GTG Val	CAG Gln	TAC Tyr	CGG	ACT Thr	GAC Asp:

	580 590	600 * * *	610	620.	
	TGG GAC CAC AGC TGG ACTrp Asp His Ser Trp Th	T GAA CAA TCA GTG r Glu Gln Ser Val	GAT TAT AGA CA Asp Tyr Arg H	AT AAG TTC is Lys Phe>	
	630 640	650 * *	. 660 * *	670 * *	
	TCC TTG CCT AGT GTG GA	AT GGG CAG AAA CGG Sp Gly Gln Lys Ar	TAC ACG TTT C Tyr Thr Phe A	GT GTT CGG rg Val Arg>	
	680 65 * *	* *	710	720	
	AGC CGC TTT AAC CCA C' Ser Arg Phe Asn Pro L	TC TGT GGA AGT GC eu Cys Gly Ser Al	T CAG CAT TGG A a Gln His Trp S	GT GAA TGG Ser Glu Trp>	
	730	740 75 * *	0 760) <u> </u>	
	AGC CAC CCA ATC CAC T Ser His Pro Ile His T	GG GGG AGC AAT AC	T TCA AAA GAG Aar Ser Lys Glu	AAC GCG TCG Asn Ala Ser>	
	770 780	790	* * *	810	•
· .	TCT GGG AAC ATG AAG C Ser Gly Asn Met Lys \	TC CTG CAG GAG CO	CC ACC TGC GTC ro Thr Cys Val	TCC GAC TAC Ser Asp Tyr>	· .
•	820 830	840	850 * * *	860 *	
	ATG AGC ATC TCT ACT Met Ser Ile Ser Thr	TGC GAG TGG AAG A Cys Glu Trp.Lys M	TG AAT GGT CCC et Asn Gly Pro	ACC AAT TGC Thr Asn Cys>	
·	870 88	0 890	900 * *	910 * *	
	AGC ACC GAG CTC CGC Ser Thr Glu Leu Arg	CTG TTG TAC CAG C Leu Leu Tyr Gln I	TG GTT TTT CTG Leu Val Phe Leu	CTC TCC GAA Leu Ser Glu>	•
	920	930 940	950	960 * *	
•	GCC CAC ACG TGT ATC Ala His Thr Cys Ile	CCT GAG AAC AAC O	GGA GGC GCG GGG Gly Gly Ala Gly	TGC GTG TGC Cys Val Cys>	
	970	980	990 * *	* *	•
	CAC CTG CTC ATG GAT His Leu Leu Met Asp	GAC GTG GTC AGT Asp Val Val Ser	GCG GAT AAC TAT Ala Asp Asn Tyt	TACA CTG GAC Thr Leu Asp>	
	1010 1020	1030	1040	1050	
•	CTG TGG GCT GGG CAG Leu Trp Ala Gly Glr	CAG CTG CTG TGG	AAG GGC TCC TT Lys Gly Ser Ph	C AAG CCC AGC e Lys Pro Ser>	
•	1060 1070	1080	1090 * *	1100 * *	
,	GAG CAT GTG AAA CCC Glu His Val Lys Pro	C AGG GCC CCA GGA D Arg Ala Pro Gly	AAC CTG ACA GT Asn Leu Thr Va	T CAC ACC AAT I His Thr Asn>	
	1110 1	120 * 1130	* 1140	1150 * *	
	GTC TCC GAC ACT CT Val Ser Asp Thr Le	G CTG CTG ACC TGG u Leu Leu Thr Trp	AGC AAC CCG TA	AT CCC CCT GAC yr Pro Pro Asp>	
•	1160	1170	80 119	1200	
•	-				
	,	•			
•••		.· ·			

AAT TAC CTG TAT AAT CAT CTC ACC TAT GCA GTC AAC ATT TGG AGT GAA Asn Tyr Leu Tyr Asn His Leu Thr Tyr Ala Val Asn Ile Trp Ser Glu> AAC GAC CCG GCA GAT TTC AGA ATC TAT AAC GTG ACC TAC CTA GAA CCC Asn Asp Pro Ala Asp Phe Arg Ile Tyr Asn Val Thr Tyr Leu Glu Pro> TCC CTC CGC ATC GCA GCC AGC ACC CTG AAG TCT GGG ATT TCC TAC AGG Ser Leu Arg Ile Ala Ala Ser Thr Leu Lys Ser Gly Ile Ser Tyr Arg> GCA CGG GTG AGG GCC TGG GCT CAG TGC TAT AAC ACC ACC TGG AGT GAG Ala Arg Val Arg Ala Trp Ala Gln Cys Tyr Asn Thr Trp Ser Glu> TGG AGC CCC AGC ACC AAG TGG CAC AAC TCC TAC AGG GAG CCC TTC GAG Trp Ser Pro Ser Thr Lys Trp His Asn Ser Tyr Arg Glu Pro Phe Glu> CAG TCC GGA GAC AAA ACT CAC ACA TGC CCA CCG TGC CCA GCA CCT GAA Gln Ser Gly Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu> CTC CTG GGG GGA CCG TCA GTC TTC CTC TTC CCC CCA AAA CCC AAG GAC Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp> ACC CTC ATG ATC TCC CGG ACC CCT GAG GTC ACA TGC GTG GTG GAC Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp> . 1570 GTG AGC CAC GAA GAC CCT GAG GTC AAG TTC AAC TGG TAC GTG GAC GGC Val Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly> GTG GAG GTG CAT AAT GCC AAG ACA AAG CCG CGG GAG GAG CAG TAC AAC Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn> AGC ACG TAC CGT GTG GTC AGC GTC CTC ACC GTC CTG CAC CAG GAC TGG Ser. Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp> . 1690 CTG AAT GGC AAG GAG TAC AAG TGC AAG GTC TCC AAC AAA GCC CTC CCA Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro> 1730· GCC CCC ATC GAG AAA ACC ATC TCC AAA GCC AAA GGG CAG CCC CGA GAA Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu>

Figure 21D

1780	,	1790		1800		*	181	O *	*	1.8	20	
CCA CAG Pro Gln	GTG TA	C ACC (ርጥር ርር	C CCA	TCC	CGG Arg	GAG (GAG . Glu	ATG Met	ACC Thr	AAG Lys	AAC Asn>
1830	ł	184		1:1			1				187	
	*	t		*	*		*	*		*		*
CAG GTC	AGC CT	rg ACC	TGC CI	G GTC	AAA	GGÇ	TTC	TAT	CCC	AGC	GAC	ATC
Gln Val	Ser Le	eu Thr	Cys Le	eu Val	Lys	Gly	Phe	Tyr	Pro	Ser	Asp	Ile>
	0.00	1	.890		19	0.0	•	is	10			1920
1	880		.09U *	*		*	*		*		*	*
GCC GTG	~ ~ ~ ~ ~ ~		NCC N	<u>አ</u> ጥ ርርር	CAG	CCG	GAG	AAC	AAC	TAC	AAG	ACC
GCC GTG Ala Val	GAG TO	CG GAG	Sor A	sn Glv	Gln	Pro	Glu	Asn	Asn	Tyr	Lys	Thr>
Ala Val	GIU I	Th Gra	Der W	J., 023	•	-						
•	1930		194	0		1950	l <u>:</u>		19	60		
*	*	*		*	*	*		*		*	*	
ACG CCT	י רככ פ	TG CTG	GAC T	CC GAG	GGC	TCC	TTC	TTC	CTC	TAT	AGC	AAG
Thr Pro	Pro V	al Leu	Asp S	er Asp	o Gly	/ Ser	Phe	Phe	Leu	Туг	Ser	Lys>
			_					•				
1970	19	080		1990			2000			2010		
•	*	*	*	*		k	*	_	* .			*
CTC AC	C GTG C	SAC AAG	AGC A	GG TG	G CA	G CA	G GGG	AAC	GTC	TTC	C TCA	A TGC
Leu Th	r Val A	Asp Lys	Ser A	rg Tr	p Gl	n Gl	n Gly	, Asr	ı Val	l Phe	e Sei	c Cys>
		2020		204	0		2.0)50			2060	
2020		2030		<i>L</i> ∪4	*	*		*		*	*	
*	* 	CAT GAG		 ኮጥር ር໓	C AA	C CA	C TA	CAC	G CA	G AA	G AG	C CTC
TCC GT	G ATG	CAT GAG His Glo	, Ala	Leu Hi	s As	n Hi	s Ty:	r Th	r Gl	n Ly	s Se	r Leu>
Ser va	T Wer	UTS OT	u Alu				_			•		
207	20	2.	080									
*	*	*	*	*								
TCC CT	rg tct	CCG GG	AAA T	TGA								•
Ser Le	eu Ser	Pro Gl	y Lys	***>			•				,	
•												

Figure 22A

	10		20	30	*	40	*
* ATG GTG	*	*	*	*. * * TO TOO	·	ጥጥሮ ሮጥG	
Met Val	Lys Pro	Ser Leu	Pro Phe	Thr Ser	Leu Leu	Phe Leu	Gln Leu>
50	£	*	70	*	80 *	90 * *	*
 בכב ביים	CTG GGA	GTG GGG Val Gly	CTG AAC Leu Asn	ACG ACA	ATT CTG	ACG CCC Thr Pro	AAT GGG Asn Gly>
100		110	120) . ••	130	*	140
* AAT GAA Asp Glu	CAC ACC	* ACA GCT Thr Ala	* * * GAT TTC Asp Phe	TTC CTC	ACC ACT	ATG CCC	ACT GAC Thr Asp>
150	,	160	• • • • • • • • • • • • • • • • • • •	170	180)	190 .
* * 'mcc cmc	* · AGT GTT	* TCC ACT	* CTG CC	* C CTC CC	* GAG GT	r CAG TG	r TTT GTG
Ser Leu	Ser Val	Ser Thi	r Leu Pro		o Giu va.		s Phe Val>
4	200	*	* *	220	*	230	240 * *
TTC AAT	GTC GAG	G TAC AT	G AAT TG t Asn Cy	C ACT TG	G AAC AG p Asn Se	C AGC TC r Ser Se	T GAG CCC r Glu Pro>
	250		260	· · · · · · · · · · · · · · · · · · ·		280	<u>.</u> .
CAG CCT Gln Pro	* T ACC AA Thr As	* C CTC AC n Leu Th	ጥ ሮጥና ርጀ	TAT TO	G TAC AA	G AAC TO	G GAT AAT er Asp Asn>
		-	310	*		*	30 · · ·
O300 333	* A GTC CA s Val Gl	ር አአር ጥ	C AGC C	AC TAT C'	TA TTC TO	CT GAA G er Glu G	AA ATC ACT lu Ile Thr>
340		350	3	60 *	370		380
* TCT GG Ser Gl	o mem e:	ላር ጥጥር ር	A AAA A	AG GAG A	TC CAC C	TC TAC C eu Tyr G	AA ACA TTT In Thr Phe>
. 39	0	400		410	4	120	, 430 * *
* GTT GT Val Va	nm	ጥሮ ሮልፎ (AC CCA C	GG GAA	CCC AGG A	AGA CAG	SCC ACA CAG
	440	4	150	460	Ç * *	470 *	480 * *
* ATG C' Met L	* TA AAA (eu Lys I	TG CAG	AAT CTG (Asn Leu	GTG ATC (CCC TGG (Pro Trp	GCT CCA (Ala Pro	GAG AAC CTA Glu Àsn Leu>
	49	0 .	500	•	510	52	0 * *
ACA C Thr L	* TT CAC Leu His	* AAA CTG Lys Leu	AGT GAA Ser Glu	TCC CAG Ser Gln	CTA GAA Leu Glu	CTG AAC Leu Asn	TGG AAC AAC Trp Asn Asn
530	A	540	55	50	560	*	570 .
AGA T Arg I	TTC TTG Phe Leu	AAC CAC Asn His.	TGT TTG Cys Leu	GAG CAC Glu His	TTG GTG Leu Val	CAG TAC Gln Tyr	CGG ACT GAC

58	0			59	90			600		*	6	10		*	62	20		
TGG Trp	* GAC Asp	CAC	: AG : Se	C 7	rGG Trp	ACT Thr	GAA Glu	CAÄ	TCA	GTC Val	GAT Asp	TA Ty	T AC	BA C cg H	AT I	AAG Lys	TTC Phe	>
	630				6	40			650		*	66	*			67	70 *	
* TCC Ser	TTG Leu	CCT Pro	* DAC Se	T (GTG Val	.* GAT Asp	GGG Gly	CAG	AA n Lys	A CGG	TA	C AC	G T'	TT C	GT Arg	GTT Val	CGG	} }
		680			,	690		.	•	700		*	71	0		÷	720) *
* AGC Ser	CGC Arg	* TT Ph	T Al	AC	* CCA Pro	CTC	TGI Cys	GG GI	A AG y Se	T GC r Al	T CA a Gl	G C	AT T is T	GG /	AGT Ser	GAA Glu	TGO	3
			730				740			75	0		ı	76	0			
AGC Ser	* CAC	C CC	. * A A o I	ጥሮ	CAC His	TG(GGGGGG	G AG	* C AA er As	T AC	x TTTO	CA A	AA C ys C	SAG Slu	AAC Asn	GGG Gly	AA As	C n>
770		•	7	80				790			80) .		ı.	810		•	
* ATG Met	AA(* G G1 s Va	rc c	* TG: Leu	CA(* G GA n Gl	G CC u Pr	* C A(o T)	CC Ť(nr Cy	* SC G! ys Va	rc To	cc c er A	SAC (rac Tyr	ATG Met	AGC Sei	AT c Il	°C .e>
8	20				830				40			850				860		
TCT Set	* C AC Th	ጥ ጥ(* GC (ys (ZAG	· ጥG	G AA	G AT	G A	* AT G sn G	GT C	CC A	CC A	TAA	TGC	AGC Ser	ACC	C G! r G:	AG lu>
	87		•			880			89	0			900				910	
* CT	c cc	* *	ጥር ' '	* TTC Lev	ፈ ጥ <i>፣</i>	* AC C? Yr G]	G C	rg G	TT T	TT C	rTG C	TC '	TCC	GAA	* GC(Ala	C CA	.C A .s T	CG hr>
ЦС	u 111		20	٠.									ç					60
ma	*		*		*		*		* GGC (. 1	k		GTG					* CTC
Сy	s I	le I	ero?	Gl	u A	sn A	sn G	ly (Gly A	Ala (3ly	Cys	Val	Cys	s Hi	s Le	eu I	eu>
			97	70		*		0 *		k			*	10	000		*	
ГА ЭМ	ng g et A	* AT (sp :	GAC Asp	· GT Va	'G G	TC A	GT G	CG	GAT Asp	AAC	TAT	ACA Thr	CTG Leu	GA(C CI	ig Ta eu T	GG (rp)	GCT Ala>
1010				102	20		*	103	0	*	10	40		*	109	50 *		*
G	* GG C ly G	יאכי	* CAG Gln	C1 L€	rg c	erg 1	rgg A	AAG	GGC Gly	TCC	TTC Phe	AAG Lys	CCC	AG Se	C G	AG C lu H	AT lis	GTG Val>
	1060)			107	70			.080		*	10	9'0		*	110) 0 *	
A L	AA (ys !	CCC Pro	AGG Arg	; G(CC (* CCA (Pro (GGA	* AAC Asn	CTG Leu	ACA Thr	GTT	CAC His	ACC Thi	C AA	AT G	TC 7	rcc Ser	GAC Asp
	-	110				112	0		1	130	•	*	114	0		*	119	50
* 4 T	(CT)	* CTG Leu	CT(G C	* :TG .eu	ACC Thr	TGG Trp	AGC Ser	AAC Asn	CCG Pro	ТАТ Туг	CCC	C CC D Pr	T GA	AC A sp A	AT '	TAC Tyr	CTG Leu
		1	160			1	170		•	11	8'0			119	0	•		1200

TAT AAT CAT CTC ACC TAT GCA GTC AAC ATT TGG AGT GAA AAC GAC CCG Tyr Asn His Leu Thr Tyr Ala Val Asn Ile Trp Ser Glu Asn Asp Pro> GCA GAT TTC AGA ATC TAT AAC GTG ACC TAC CTA GAA CCC TCC CTC CGC Ala Asp Phe Arg Ile Tyr Asn Val Thr Tyr Leu Glu Pro Ser Leu Arg> ATC GCA GCC AGC ACC CTG AAG TCT GGG ATT TCC TAC AGG GCA CGG GTG Ile Ala Ala Ser Thr Leu Lys Ser Gly Ile Ser Tyr Arg Ala Arg Val> AGG GCC TGG GCT CAG AGC TAT AAC ACC ACC TGG AGT GAG TGG AGC CCC Arg Ala Trp Ala Gln Ser Tyr Asn Thr Trp Ser Glu Trp Ser Pro> AGC ACC AAG TGG CAC AAC TCC TAC AGG GAG CCC TTC GAG CAG TCC GGA Ser Thr Lys Trp His Asn Ser Tyr Arg Glu Pro Phe Glu Gln Ser Gly> GAC AAA ACT CAC ACA TGC CCA CCG TGC CCA GCA CCT GAA CTC CTG GGG Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly> GGA CCG TCA GTC TTC CTC TTC CCC CCA AAA CCC AAG GAC ACC CTC ATG Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu Met> ATC TCC CGG ACC CCT GAG GTC ACA TGC GTG GTG GTG GAC GTG AGC CAC Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser His> GAA GAC CCT GAG GTC AAG TTC AAC TGG TAC GTG GAC GGC GTG GAG GTG Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val Glu Val> CAT AAT GCC AAG ACA AAG CCG CGG GAG GAG CAG TAC AAC AGC ACG TAC His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr Tyr> CGT GTG GTC AGC GTC CTC ACC GTC CTG CAC CAG GAC TGG CTG AAT GGC Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn Gly> AAG GAG TAC AAG TGC AAG GTC TCC AAC AAA GCC CTC CCA GCC CCC ATC Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro Ile> GAG AAA ACC ATC TCC AAA GCC AAA GGG CAG CCC CGA GAA CCA CAG GTG Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln Val>

Pro Gly Lys ***>

TAC ACC CTG CCC CCA TCC CGG GAT GAG CTG ACC AAG AAC CAG GTC AGC. Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val Ser> CTG ACC TGC CTG GTC AAA GGC TTC TAT CCC AGC GAC ATC GCC GTG GAG Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val Glu> TGG GAG AGC AAT GGG CAG CCG GAG AAC AAC TAC AAG ACC ACG CCT CCC Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro Pro> GTG CTG GAC TCC GAC GGC TCC TTC TTC CTC TAT AGC AAG CTC ACC GTG Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr Val> GAC AAG AGC AGG TGG CAG CAG GGG AAC GTC TTC TCA TGC TCC GTG ATG Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val Met> CAT GAG GCT CTG CAC AAC CAC TAC ACG CAG AAG AGC CTC TCC CTG TCT His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu Ser> CCG GGT AAA TGA

Figure 23A

		•		
10	20	30	40	
* * * * ATG GTG AAG CCA TCA TTA	* * *	* ጉል ጥርር ርጥሮ	* * * TTA TTC CTG	* CAG CTG
Met Val Lys Pro Ser Leu	Pro Phe T	hr Ser Leu	Leu Phe Leu	Gln Leu>
	70	80	90	
50 60 . * * *	*	* *	* *	*
CCC CTG CTG GGA GTG GGG	CTG AAC A	CG ACA ATT	CTG ACG CCC	AAT GGG Asn Glv>
Pro Leu Leu Gly Val Gly	Leu Asn 1			
100 110	120	*	.30 * *	140
AAT GAA GAC ACC ACA GCT	ርልጥ ጥጥር ባ	TTC CTG ACC	ACT ATG CCC	ACT GAC
AAT GAA GAC ACC ACA GC. Asn Glu Asp Thr Thr Ala	Asp Phe I	Phe Leu Thi	Thr Met Pro	nr Asp>
150 160	1.	70	180	190
* * * * * TCC CTC AGT GTT TCC AC	* ኮ ሮሞG ሮሮሮ (* * CTC CCA GA		r TTT GTG
Ser Leu Ser Val Ser Th	r Leu Pro	Leu Pro Gl	u Val Gln Cy	s Phe Val>
0.4		220	230	240
	* *	* * *	* * *	* · *
TTC AAT GTC GAG TAC AT Phe Asn Val Glu Tyr Me	G AAT TGC	ACT TGG AA Thr Trp As	n Ser Ser Se	r Glu Pro>
pne Ash val Glu 171			280	
250	2 <u>6</u> 0	270 * *	* *	*
CAG CCT ACC AAC CTC AC	OT CTG CAT	TAT TGG TA	AC AAG AAC TO	CG GAT AAT er Asp Asp>
Gln Pro Thr Asn Leu T	ir Leu His	TAI TIP I		
290 300	310 * *	*	0	30 * *
GAT AAA GTC CAG AAG T	ככ אכר כאר	TAT CTA T	TC TCT GAA G	AA ATC ACT
GAT AAA GTC CAG AAG T Asp Lys Val Gln Lys C	ys Ser His	Tyr Leu P	he Ser Glu G	lu lie Thr>
340 350	360)	370	380
* * * *	* * אא אא אמי	·	AC CTC TAC C	AA ACA TTT
Ser Gly Cys Gln Leu (Sln Lys Lys	Glu Ile F	lis Leu Tyr (In Thr Phe>
		410	420	430
3,50	* *	*	* * . NGC NGN CNG !	* * CCC ACA CAG
GTT GTT CAG CTC CAG Val Val Gln Leu Gln	GAC CCA CG Asp Pro Ar	g GAA CCC . g Glu Pro .	Arg Arg Gln	Ala Thr Gln>
		460 ·	470	480
440 * * *	450 * *	*	* *	* *
ATG CTA AAA CTG CAG Met Leu Lys Leu Gln	AAT CTG GT	G ATC CCC	TGG GCT CCA Trp Ala Pro	GAG AAC CTA Glu Asn Leu>
Met Leu Lys Leu Gin	Wall Dea 40			
490	500 *	* * *	*	20 * *
ACA CTT CAC AAA CTG	AGT GAA TO	CC CAG CTA	GAA CTG AAC	TGG AAC AAC
Thr Leu His Lys Leu	Ser Glu S	er din ren	Gid bed Asii	11p 11011 11011
530 540	550	*	560	570 * *
AGA TTC TTG AAC CAC	TGT TTG G	AG CAC TTG	GTG. CAG TAC	CGG ACT GAC
Arg Phe Leu Asn His	Cys Leu G	lu His Leu	Val Gln Tyr	Arg Thr Asp

32/74 Figure 23B

TGG GAC CAC AGC TGG ACT GAA CAA TCA GTG GAT TAT AGA CAT AAG TTC Trp Asp His Ser Trp Thr Glu Gln Ser Val Asp Tyr Arg His Lys Phe> TCC TTG CCT AGT GTG GAT GGG CAG AAA CGC TAC ACG TTT CGT GTT CGG Ser Leu Pro Ser Val Asp Gly Gln Lys Arg Tyr Thr Phe Arg Val Arg> AGC CGC TTT AAC CCA CTC TGT GGA AGT GCT CAG CAT TGG AGT GAA TGG Ser Arg Phe Asn Pro Leu Cys Gly Ser Ala Gln His Trp Ser Glu Trp> AGC CAC CCA ATC CAC TGG GGG AGC AAT ACT TCA AAA GAG AAC GCG TCG Ser His Pro Ile His Trp Gly Ser Asn Thr Ser Lys Glu Asn Ala Ser> TCT GGG AAC ATG AAG GTC CTG CAG GAG CCC ACC TGC GTC TCC GAC TAC Ser Gly Asn Met Lys Val Leu Gln Glu Pro Thr Cys Val Ser Asp Tyr> ATG AGC ATC TCT ACT TGC GAG TGG AAG ATG AAT GGT CCC ACC AAT TGC Met Ser Ile Ser Thr Cys Glu Trp Lys Met Asn Gly Pro Thr Asn Cys> AGC ACC GAG CTC CGC CTG TTG TAC CAG CTG GTT TTT CTG CTC TCC GAA Ser Thr Glu Leu Arg Leu Leu Tyr Gln Leu Val Phe Leu Leu Ser Glu> GCC CAC ACG TGT ATC CCT GAG AAC AAC GGA GGC GCG GGG TGC GTG TGC Ala His Thr Cys Ile Pro Glu Asn Asn Gly Gly Ala Gly Cys Val Cys> CAC CTG CTC ATG GAT GAC GTG GTC AGT GCG GAT AAC TAT ACA CTG GAC His Leu Leu Met Asp Asp Val Val Ser Ala Asp Asn Tyr Thr Leu Asp> CTG TGG GCT GGG CAG CAG CTG CTG TGG AAG GGC TCC TTC AAG CCC AGC Leu Trp Ala Gly Gln Gln Leu Leu Trp Lys Gly Ser Phe Lys Pro Ser> GAG CAT GTG AAA CCC AGG GCC CCA GGA AAC CTG ACA GTT CAC ACC AAT Glu His Val Lys Pro Arg Ala Pro Gly Asn Leu Thr Val His Thr Asn> GTC TCC GAC ACT CTG CTG CTG ACC TGG AGC AAC CCG TAT CCC CCT GAC Val Ser Asp Thr Leu Leu Leu Thr Trp Ser Asn Pro Tyr Pro Pro Asp>

AAT TAC CTG TAT AAT CAT CTC ACC TAT GCA GTC AAC ATT TGG AGT GAA Asn Tyr Leu Tyr Asn His Leu Thr Tyr Ala Val Asn Ile Trp Ser Glu> AAC GAC CCG GCA GAT TTC AGA ATC TAT AAC GTG ACC TAC CTA GAA CCC Asn Asp Pro Ala Asp Phe Arg Ile Tyr Asn Val Thr Tyr Leu Glu Pro> TCC CTC CGC ATC GCA GCC AGC ACC CTG AAG TCT GGG ATT TCC TAC AGG Ser Leu Arg Ile Ala Ala Ser Thr Leu Lys Ser Gly Ile Ser Tyr Arg> GCA CGG GTG AGG GCC TGG GCT CAG AGC TAT AAC ACC ACC TGG AGT GAG Ala Arg Val Arg Ala Trp Ala Gln Ser Tyr Asn Thr Thr Trp Ser Glu> TGG AGC CCC AGC ACC AAG TGG CAC AAC TCC TAC AGG GAG CCC TTC GAG Trp Ser Pro Ser Thr Lys Trp His Asn Ser Tyr Arg Glu Pro Phe Glu> 1440. CAG TCC GGA GAC AAA ACT CAC ACA TGC CCA CCG TGC CCA GCA CCT GAA Gln Ser Gly Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu> CTC CTG GGG GGA CCG TCA GTC TTC CTC TTC CCC CCA AAA CCC AAG GAC Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp> ACC CTC ATG ATC TCC CGG ACC CCT GAG GTC ACA TGC GTG GTG GAC Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp> GTG AGC CAC GAA GAC CCT GAG GTC AAG TTC AAC TGG TAC GTG GAC GGC Val Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly> GTG GAG GTG CAT AAT GCC AAG ACA AAG CCG CGG GAG GAG CAG TAC AAC Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn> AGC ACG TAC CGT GTG GTC AGC GTC CTC ACC GTC CTG CAC CAG GAC TGG Ser Thr Tyr Arg Val Val Ser Val Leu Thr. Val Leu His Gln Asp Trp> CTG AAT GGC AAG GAG TAC AAG TGC AAG GTC TCC AAC AAA GCC CTC CCA Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro> GCC CCC ATC GAG AAA ACC ATC TCC AAA GCC AAA GGG CAG CCC CGA GAA

Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu>

Figure 23D

CCA CAG GTG TAC ACC CTG CCC CCA TCC CGG GAT GAG CTG ACC AAG AAC Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn> CAG GTC AGC CTG ACC TGC CTG GTC AAA GGC TTC TAT CCC AGC GAC ATC Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile> GCC GTG GAG TGG GAG AGC AAT GGG CAG CCG GAG AAC AAC TAC AAG ACC Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr> ACG CCT CCC GTG CTG GAC TCC GAC GGC TCC TTC TTC CTC TAT AGC AAG Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys> CTC ACC GTG GAC AAG AGC AGG TGG CAG CAG GGG AAC GTC TTC TCA TGC Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys> TCC GTG ATG CAT GAG GCT CTG CAC AAC CAC TAC ACG CAG AAG AGC CTC Ser Val Met His Glu Ala, Leu His Asn His Tyr Thr Gln Lys Ser Leu> TCC CTG TCT CCG GGT AAA TGA Ser Leu Ser Pro Gly Lys ***>

	•			
10	20	30	40	*
* * * . *	* *		፣ ኮሬ ርፑሬ GCC (CG CCG
* * * * * ATG GTG GCC GTC GGC Met Val Ala Val Gly	TGC GCG CTG C	eu Ala Ala Lo	eu Leu Ala A	Ala Pro>
Met Val Ala Val Gly	cys Ara bed -			•
50 60	70	80	90	•
	* *	* * * *	* * አር ርኔር ርጥር !	CCA AGA
* * * * GGA GCG GCG GCC Gly Ala Ala Leu Ala	CCA AGG CGC	rge pro Ala G	in Glu Val	Ala Arg>
Gly Ala Ala Leu Ala	Pro Arg Arg	cys iio iii -		_
100 110	120	130) 1	.40 .
100	* *	*	* * ·	* . TCC CCC
GGC GTG CTG ACC AG	CTG CCA GGA	GAC AGC GIG A	Thr Leu Thr	Cys Pro>
GGC GTG CTG ACC AGG Gly Val Leu Thr Sex	c Leu Pro Gly	Asp ser var		- 4
150	160	.70	180	190
4.50		* *	* *	*
* * * * GGG GTA GAG CCG GA	A GAC AAT GCC	ACT GTT CAC	TGG GTG CTC Trn Val Leu	Ara Lvs>
GGG GTA GAG CCG GA Gly Val Glu Pro Gl	u Asp Asn Ala	Thr val his	TIP VAI LCA	
200	210	220	230	240
200	*	* *	*	* *
CCG GCT GCA GGC TC	C CAC CCC AGC	AGA TGG GCT	GGC ATG GGA	AGG AGG
CCG GCT GCA GGC TC Pro Ala Ala Gly Se	er His Pro Ser	Arg Trp Ala	GIA WEL GIA	Aly Aly
•	260	270	280	
250 * *	. •	* *	* *	*
	CG GTG CAG CTC	CAC GAC TCT	GGA AAC TAT	r TCA TGC
CTG CTG CTG AGG To Leu Leu Leu Arg S	er Val Gln Le	His Asp Ser	GIY Asn Ty	r ser cys>
	310	320	33	
290 300		* *	*	* *
* * * * TAC CGG GCC GGC C	GC CCA GCT GG	G ACT GTG CAC	TTG CTG GT	G GAT GTT
TAC CGG GCC GGC C Tyr Arg Ala Gly A	rg Pro Ala Gl	y Thr Val His	s Leu Leu va	II Wah Agra
	2.0	-	370	380
740		* *	* *	*
* * CCC CCC GAG GAG	CCC CAG CTC TO	C TGC TTC CGG	G AAG AGC CO	CC CTC AGC
CCC CCC GAG GAG (Pro Pro Glu Glu	Pro Gln Leu Se	er Cys Phe Ar	g Lys Ser P	to rea serv
	•	410	420	430
390	400	* *	*	* *
K	GAG TGG GGT C	CT CGG AGC AC	C CCA TCC C	TG ACG ACA
AAT GTT GTT TGT Asn Val Val Cys	Glu Trp Gly P	ro Arg Ser Th	ir Pro Ser L	eu mi mi
		460	470	480
440	450 * *	* *	* *	* *
* * AAG GCT GTG CTC	TTG GTG AGG A	AG TTT CAG A	AC AGT CCG	SCC GAA GAC
Lys Ala Val Leu	Leu Val Arg I	ys Phe Gln A	sn Ser Pro I	Ala Glu Asp
	500	510	52	
490		* *	*	* *
* * TTC CAG GAG CCG	TGC CAG TAT	TCC CAG GAG T	CC CAG AAG	TTC TCC TGC
TTC CAG GAG CCG Phe Gln Glu Pro	Cys Gln Tyr	Ser Gln Glu S	ser Gin Lys	Aug Ser Chas
F 4.0	.		50	570
530 540		* *	* *	* *
CAG TTA GCA GTO	CCG GAG GGA	GAC AGC TCT	TTC TAC ATA	GTG TCC ATG
The Walter	Pro Glu GlV	Asp Ser Ser	Phe Tyr Ile	val Ser Met

	580 59	600	* * *	620 *
	TGC GTC GCC AGT A	AGT GTC GGG AGC AAG Ser Val Gly Ser Lys	TTC AGC AAA ACT CAA	ACC TTT Thr Phe>
	630.	640 650	660 * *	670
	CAG GGT TGT GGA Gln Gly Cys Gly	ATC TTG CAG CCT GAT Ile Leu Gln Pro Asp	CCG CCT GCC AAC ATO	C ACA GTC Thr Val>
	680	690	700 710	720
	* * ACT GCC GTG GCC Thr Ala Val Ala	AGA AAC CCC CGC TGC Arg Asn Pro Arg Tr	CTC AGT GTC ACC TG Leu Ser Val Thr Tr	G CAA GAC p Gln Asp>
	730	740	750	*
	CCC CAC TCC TGG Pro His Ser Trp	AAC TCA TCT TTC TA Asn Ser Ser Phe Ty	C AGA CTA CGG TTT GA r Arg Leu Arg Phe Gl	G CTC AGA u Leu Arg>
	770 780	790 * *	* * * *	.0 * *
	TAT CGG GCT GAA Tyr Arg Ala Glu	CGG TCA AAG ACA TT Arg Ser Lys Thr Ph	C ACA ACA TGG ATG G' he Thr Thr Trp Met Va	CC AAG GAC al Lys Asp>
	820	830 840	850 * * *	860 *
•	CTC CAG CAT CAC	TGT GTC ATC CAC G	AC GCC TGG AGC GGC C sp Ala Trp Ser Gly L	TG AGG CAC eu Arg His>
·	870	880 89	0 900 * * *	910 * *
	GTG GTG CAG CTT	r CGT GCC CAG GAG G	AG TTC GGG CAA GGC Clu Phe Gly Gln Gly C	AG TGG AGC lu Trp Ser>
	920	930	940 950	960
	* * *	* * * * G GAG GCC ATG GGC A O Glu Ala Met Gly T	CG CCT TGG ACA GAA 'Chr Pro Trp Thr Glu	* * CC AGG AGT Ser Arg Ser>
	970	980	990 100	•
· · · · · · · · · · · · · · · · · · ·	* * CCT CCA GCT GA Pro Pro Ala Gl	* G AAC GAG GTG TCC A u Asn Glu Val Ser '	ACC CCC ATG ACC GGT Thr Pro Met Thr Gly	GGC GCG CCT Gly Ala Pro>
•	1010 102	1030	1040 1	050
	* * TCA GGT GCT CA Ser Gly Ala G	AG CTG GAA CTT CTA In Leu Glu Leu Leu	GAC CCA TGT GGT TAT Asp Pro Cys Gly Tyr	ATC AGT CCT Ile Ser Pro>
	1060	1070 * 1080	* 1090 * * *	1100
· .	GAA TCT CCA G Glu Ser Pro V	TT GTA CAA CTT CAT al Val Gln Leu His	TCT AAT TTC ACT GCA Ser Asn Phe Thr Ala	GTT TGT GTG Val Cys Val>
· · .	1110	* * *	1140	1150 * *
	CTA AAG GAA A Leu Lys Glu L	AA TGT ATG GAT TAT Lys Cys Met Asp Tyr	TTT CAT GTA AAT GCT Phe His Val Asn Ala	AAT TAC ATT Asn Tyr Ile>
	1160	1170	1180 1190	1200
	,		•	
	•			•

	GTC TGG AAA ACA AAC CAT TTT ACT ATT CCT AAG GAG CAA TAT ACT ATC Val Trp Lys Thr Asn His Phe Thr Ile Pro Lys Glu Gln Tyr Thr Ile>
	1210 1220 1230 1240
	ATA AAC AGA ACA GCA TCC AGT GTC ACC TTT ACA GAT ATA GCT TCA TTA Ile Asn Arg Thr Ala Ser Ser Val Thr Phe Thr Asp Ile Ala Ser Leu>
•	1250 1260 1270 1280 1290
	AAT ATT CAG CTC ACT TGC AAC ATT CTT ACA TTC GGA CAG CTT GAA CAG Asn Ile Gln Leu Thr Cys Asn Ile Leu Thr Phe Gly Gln Leu Glu Gln>
	1300 1310 1320 1330 1340
•	AAT GTT TAT GGA ATC ACA ATA ATT TCA GGC TTG CCT CCA GAA AAA CCT Asn Val Tyr Gly Ile Thr Ile Ile Ser Gly Leu Pro Pro Glu Lys Pro>
	1350 1360 1370 1380 1390
	AAA AAT TTG AGT TGC ATT GTG AAC GAG GGG AAG AAA ATG AGG TGT GAG Lys Asn Leu Ser Cys Ile Val Asn Glu Gly Lys Lys Met Arg Cys Glu>
	1400 1410 1420 1430 1440
	* TGG GAT GGT GGA AGG GAA ACA CAC TTG GAG ACA AAC TTC ACT TTA AAA Trp Asp Gly Gly Arg Glu Thr His Leu Glu Thr Asn Phe Thr Leu Lys>
•	1450 1460 1470 1480
•	TCT GAA TGG GCA ACA CAC AAG TTT GCT GAT TGC AAA GCA AAA CGT GAC Ser Glu Trp Ala Thr His Lys Phe Ala Asp Cys Lys Ala Lys Arg Asp>
	1490 1500 1510 1520 1530
•	ACC CCC ACC TCA TGC ACT GTT GAT TAT TCT ACT GTG TAT TTT GTC AAC Thr Pro Thr Ser Cys Thr Val Asp Tyr Ser Thr Val Tyr Phe Val Asn>
	1540 1550 1560 1570 1580
	* * * * * * * * * * * * * * * * * * *
	1590 1600 1610 1620 1630
	GAT CAT ATC AAT TTT GAT CCT GTA TAT AAA GTG AAG CCC AAT CCG CCA Asp His Ile Asn Phe Asp Pro Val Tyr Lys Val Lys Pro Asn Pro Pro>
	1640 1650 1660 1670 1680
	CAT AAT TTA TCA GTG ATC AAC TCA GAG GAA CTG TCT AGT ATC TTA AAA His Asn Leu Ser Val Ile Asn Ser Glu Glu Leu Ser Ser Ile Leu Lys>
•	1690 1700 1710 1720
	TTG ACA TGG ACC AAC CCA AGT ATT AAG AGT GTT ATA ATA CTA AAA TAT Leu Thr Trp Thr Asn Pro Ser Ile Lys Ser Val Ile Ile Leu Lys Tyr>
	1730 1740 1750 1760 1770
	AAC ATT CAA TAT AGG ACC AAA GAT GCC TCA ACT TGG AGC CAG ATT CCT Asn Ile Gln Tyr Arg Thr Lys Asp Ala Ser Thr Trp Ser Gln Ile Pro

Figure 24D

		•													
1780	•	17	90		*	800		*	183	¥	*	18	320		
CCT GAA	GAC Asp	ACA Thr	GCA Ala	TCC Ser	ACC	CGA Arg	TCT Ser	TCA Ser	TTC Phe	ACT Thr	GTC Val	CAA Gln	GAC Asp	CTT Leu:	>
1830)		184	10		1	850		*	1860		*	18	70 *	
* AAA CCT	r TTT o Phe	* ACA Thr	GAA Glu	ጥልጥ	GTG	· TTT	AGG	ATT	r CGC e Arg	TGT Cys	ATG Met	AAG Lys	GAA Glu	GAT Asp	>
*	1880	٠	*	1890		*	19	000	*	1	910			1920)
GGT AAG	G GGA s Gly	TAC	ጥርር	AGT Ser	GAC Asp	TGC Trp	AG:	r GA	A GAA u Glu	GCA Ala	AGT Ser	GGG Gly	ATC Ile	ACC Thr	; ;>
	19	30-		1	940		*	195	0	•	19	960	•	r	
TAT GA Tyr Gl	A GAT u Asp	* ! AGA > Arç	* CCA Pro	TCT Sex	* AA Lys	A GC	A CC	A AG	T TT	C'TG(e'Tr	TAT TY	r AAA	ATA	GA' Asj	p>
1970		1980)	•	1	990		*	2000		•	201	0	*	
CCA TO	* CC CA' er Hi	r AC' s Th:	r CA/ r Gli	* A GGG A Gly	TA TY	C AG	A AC	T G	ra ca al Gl	A CT n Le	C GT u Va	G TG	G AA	3 AC s Th	A r>
2020			2030			204			2			*	2060		
TTG Co	CT CC ro Pr	* T TT o Ph	T GA le Gl	A GC	C AA	T GO	A A	A A	TC TI	G GA	AT TA	T GA	A GT u Va	G AC	er er>
	70		2				209		*:	210		4		2110	
* CTC A Leu T	* CA AC hr Ai	בא ייני	C AA	א ידיכ	A CA	T T	TA C	AA A	AT T	AC AC	CA G	rr Al	AT GO sn Al	C AC	CA hr>
•	2120					•		2140) .	*	215		*	21	
AAA (Lys I	CTG AC	* CA G' hr Va	מ מיזי	AT C'	 ፑር ል	CA A	AT G	AT (CGC T Arg T	AT C	TA G eu A	CA A la T	CC C' hr L	TA A eu T	CA hr>
		2170		*	218	0		2	190 *		*	2200		*	
GTA . Val	* AGA A Arg A	AT C sn L	TT G	ጥጥ G	GC A	AA 7	CA C Ser A	Asp	GCA (SCT G	TT T	TA AT.	CT A	TC C	CCT Pro>
2210		22	220		*	223) *	*	22	40 *	. ,	, 22 * ·	250 * .	•	*
CCC	TGT (יאר ח	TTT C	AA C	CT. A	ACT (CAC His	CCT Pro	GTA . Val	ATG (GAT (Asp	CTT I Leu	AAA (Lys)	3CA /	TTC Phe>
226	50	*	227	7 O *		2	280		*	229	0	*	23	00	
CCC Pro	AAA (ርልጥ :	AAC <i>l</i> Asn l	ATG (Met	CTT Leu	TGG Trp	GTG Val	GAA Glu	TGG Trp	ACT Thr	ACT Thr	CCA Pro	AGG Arg	GAA Glu	TCT Ser>
*	2310		*	232	0	*	23	330		* 2	340		*	239	50 *
GTA.	AAG Lys	AAA Lys	TAT	ATA Ile	CTT Leu	GAG Glu	TGG Trp	TGT Cys	GTG Val	TTA Leu	TCA Ser	GAT Asp	AAA Lys	GCA Ala	CCC Pro
	23	60		2	370			23	80		2:	390		i	2400

Figure 24E

* * * * * * * * * * * * * * * * * * * *	* *
TGT ATC ACA GAC TGG CAA CAA GAA GAT GGT ACC GTG CAT CGG Cys Ile Thr Asp Trp Gln Gln Glu Asp Gly Thr Val His Arc	C ACC TAT g Thr Tyr>
2410 2420 2430 2440	*
TTA AGA GGG AAC TTA GCA GAG AGC AAA TGC TAT TTG ATA AC Leu Arg Gly Asn Leu Ala Glu Ser Lys Cys Tyr Leu Ile Th	A GTT ACT r Val Thr>
2450 2460 2470 2480 249	·
CCA GTA TAT GCT GAT GGA CCA GGA AGC CCT GAA TCC ATA AF Pro Val Tyr Ala Asp Gly Pro Gly Ser Pro Glu Ser Ile Ly	AG GCA TAC ys Ala Tyr>
2500 2510 2520 2530	2540 *
CTT AAA CAA GCT CCA CCT TCC AAA GGA CCT ACT GTT CGG AG Leu Lys Gln Ala Pro Pro Ser Lys Gly Pro Thr Val Arg T	CA AAA AAA hr Lys Lys>
2550 2560 2570 2580 * * * * * * * *	2590 * *
GTA GGG AAA AAC GAA GCT GTC TTA GAG TGG GAC CAA CTT C Val Gly Lys Asn Glu Ala Val Leu Glu Trp Asp Gln Leu F	CT GTT GAT
2600 2610 2620 2630	2640 * *
GTT CAG AAT GGA TTT ATC AGA AAT TAT ACT ATA TTT TAT A	AGA ACC ATC Arg Thr Ile>.
2650 2660 2670 268	* *
ATT GGA AAT GAA ACT GCT GTG AAT GTG GAT TCT TCC CAC Ile Gly Asn Glu Thr Ala Val Asn Val Asp Ser Ser His	ACA GAA TAT Thr Glu Tyr>
2690 2700 * * * * * *	.730 * *
ACA TTG TCC TCT TTG ACT AGT GAC ACA TTG TAC ATG GTA Thr Leu Ser Ser Leu Thr Ser Asp Thr Leu Tyr Met Val	CGA ATG GCA Arg Met Ala>
2740 2750 2760 2770	2780
GCA TAC ACA GAT GAA GGT GGG AAG GAT GGT CCA GAA TTC Ala Tyr Thr Asp Glu Gly Gly Lys Asp Gly Pro Glu Phe	ACT TTT ACT Thr Phe Thr>
2790 2800 2810 2820	2830 * *
ACC CCA AAG TTT GCT CAA GGA GAA ATT GAA TCC GGG GGC Thr Pro Lys Phe Ala Gln Gly Glu Ile Glu Ser Gly Gly	GAC AAA ACT Asp Lys Thr>
2840 2850 2860 2870	* *
CAC ACA TGC CCA CCG TGC CCA GCA CCT GAA CTC CTG GGC His Thr Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly	GGA CCG TCA Gly Pro Ser>
2890 2500 * * * *	920
GTC TTC CTC TTC CCC CCA AAA CCC AAG GAC ACC CTC AT Val Phe Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu Me	G ATC TCC CGG t lle Ser Arg>
2930 2940 2950 2960	2970
ACC CCT GAG GTC ACA TGC GTG GTG GTG GAC GTG AGC CA	AC GAA GAC CCT.

* * * >

Thr Pro Glu Val Thr Cys Val Val Val Asp Val Ser His Glu Asp Pro> GAG GTC AAG TTC AAC TGG TAC GTG GAC GGC GTG GAG GTG CAT AAT GCC Glu Val Lys Phe Asn Trp Tyr Val Asp Gly Val Glu Val His Asn Ala> AAG ACA AAG CCG CGG GAG GAG CAG TAC AAC AGC ACG TAC CGT GTG GTC Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val Val> AGC GTC CTC ACC GTC CTG CAC CAG GAC TGG CTG AAT GGC AAG GAG TAC Ser Val Leu Thr Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu Tyr> AAG TGC AAG GTC TCC AAC AAA GCC CTC CCA GCC CCC ATC GAG AAA ACC Lys Cys Lys Val Ser Asn Lys Ala Leu Pro Ala Pro Ile Glu Lys Thr> ATC TCC AAA GCC AAA GGG CAG CCC CGA GAA CCA CAG GTG TAC ACC CTG Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu Pro Gln Val Tyr Thr Leu> CCC CCA TCC CGG GAT GAG CTG ACC AAG AAC CAG GTC AGC CTG ACC TGC Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn Gln Val Ser Leu Thr Cys> 3290 -CTG GTC AAA GGC TTC TAT CCC AGC GAC ATC GCC GTG GAG TGG GAG AGC Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile Ala Val Glu Trp Glu Ser> AAT GGG CAG CCG GAG AAC AAC TAC AAG ACC ACG CCT CCC GTG CTG GAC Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr Thr Pro Pro Val Leu Asp> TCC GAC GGC TCC TTC TTC CTC TAC AGC AAG CTC ACC GTG GAC AAG AGC Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys Leu Thr Val Asp Lys Ser> AGG TGG CAG CAG GGG AAC GTC TTC TCA TGC TCC GTG ATG CAT GAG GCT Arg Trp Gln Gln Gly Asn Val Phe Ser Cys Ser Val Met His Glu Ala> CTG CAC AAC CAC TAC ACG CAG AAG AGC CTC TCC CTG TCT CCG GGT AAA Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys> TGA

Figure 25A.

10 20 30 40
* * * * * * * * * * * * * * * * * * *
Met Val Ala Val Gly Cys Ala Leu Leu Ala Mac 120 200 200
50 60 70 80 90
* * * * * * GGA GCG CTG GCC CCA AGG CGC TGC CCT GCG CAG GAG GTG GCA AGA GGA GCG GCG CTG GCC CCA AGG CGC TGC CCT GCG CAG GAG GTG GCA AGA Gly Ala Ala Leu Ala Pro Arg Arg Cys Pro Ala Gln Glu Val Ala Arg>
130 130 140
* * * * * * * * * * * * * * * * * * *
GGC GTG CTG ACC AGT CTG CCA GGA GAC AGC OTO THE Leu Thr Cys Pro> Gly Val Leu Thr Ser Leu Pro Gly Asp Ser Val Thr Leu Thr Cys Pro>
150 160 170 180 190
* * * * * * * * * * * * * * * * * * *
200 210 220 230 240
* * * * * * CCG GCT GCA GGC TCC CAC CCC AGC AGA TGG GCT GGC ATG GGA AGG AGG CCG GCT GCT GCA ATG ATG ACG AGG AGG CCG GCT GCC ATG GGA AGG AGG AGG CCG GCT GCC ATG GCA AGG AGG AGG CCG GCT GCC ATG GCA AGG AGG AGG AGG AGG AGG GCT GCC ATG GCA ATG ATG ATG ATG ATG ATG ATG ATG ATG AT
250 260 270 280 **
* * * *
* * * * * CTG CTG CTG CTG CAC CTC CAC GAC TCT GGA AAC TAT TCA TGC CTG CTG CTG AGG TCG GTG CAG CTC CAC GAC TCT GGA AAC TAT TCA TGC CTG CTG CTG CTG CTG CTG CTG CTG CTG
290 300 310 320 330
* * * * * * TOTAL THE TAC CGG GCC GCC CCA GCT GGG ACT GTG CAC TTG CTG GTG GAT GTT TAC CGG GCC GGC CGC CCA GCT GGG ACT GTG CAC TTG CTG GTG GAT GTT TAC CGG GCC GGC CCA GCT GGG ACT GTG CAC TTG CTG GTG GAT GTT TAC CTG GTG GTG GAT GTT TAC CTG GTG GTG GAT GTT TAC CTG GTG GTG GAT GTT TAC CTG GTG GTG GTG GAT GTT TAC CTG GTG GTG GTG GTG GTG GTG GTG GTG GT
340 350 360 370 380
* * * * * * * * * * * * * * * * * * *
390 400 410 420 430
* * * * * * * * * * * * * * * * * * *
440 450 460 470 . 480
* * * * * * * * * * * * * * * * * * *
Lys Ala Val Leu Leu Val Arg Lys Pile Gin Asi. Con 120
490 500 510 520
* * * TTC CAG GAG CCG TGC CAG TAT TCC CAG GAG TCC CAG AAG TTC TCC TGC Phe Gln Glu Pro Cys Gln Tyr Ser Gln Glu Ser Gln Lys Phe Ser Cys>
530 540 550 560 570
* CAG TTA GCA GTC CCG GAG GGA GAC AGC TCT TTC TAC ATA GTG TCC ATG Gln Leu Ala Val Pro Glu Gly Asp Ser Ser Phe Tyr Ile Val Ser Met>

Figure 25B

		580 590 600 610 620	
		* * * * * * * * TGC GTC GCC AGT AGT GTC GGG AGC AAG TTC AGC AAA ACT CAA ACC TTT Cys Val Ala Ser Ser Val Gly Ser Lys Phe Ser Lys Thr Gln Thr Phe>	
·		630 640 650 660 670	
7		CAG GGT TGT GGA ATC TTG CAG CCT GAT CCG CCT GCC AAC ATC ACA GTC Gln Gly Cys Gly Ile Leu Gln Pro Asp Pro Pro Ala Asn Ile Thr Val>	
		680 690 700 710 720	
		* * * * * * * * * * * * * * * * * * *	
		730 740 750 760	
	· · · ·	* * * * * * * * * * * * * * * * * * *	
		770 780 790 800 810	
		* * * * * * * * * * * * * * * * * * *	
		820 830 840 850 860	
		CTC CAG CAT CAC TGT GTC ATC CAC GAC GCC TGG AGC GGC CTG AGG CAC Leu Gln His His Cys Val Ile His Asp Ala Trp Ser Gly Leu Arg His>	
		870 880 890 900 910	
		* * * * GTG CAG CTT CGT GCC CAG GAG GAG TTC GGG CAA GGC GAG TGG AGC Val Val Gln Leu Arg Ala Gln Glu Glu Phe Gly Gln Gly Glu Trp Ser>	
	•	920 930 940 950 960	
	·	GAG TGG AGC CCG GAG GCC ATG GGC ACG CCT TGG ACA GAA TCG CGA TCG Glu Trp Ser Pro Glu Ala Met Gly Thr Pro Trp Thr Glu Ser Arg Ser>	
		970 980 990 1000	
		CCT CCA GCT GAG AAC GAG GTG TCC ACC CCC ATG GAA CTT CTA GAC CCA Pro Pro Ala Glu Asn Glu Val Ser Thr Pro Met Glu Leu Asp Pro>	
	•	1010 1020 1030 1040 1050	
		TGT GGT TAT ATC AGT CCT GAA TCT CCA GTT GTA CAA CTT CAT TCT AAT Cys Gly Tyr Ile Ser Pro Glu Ser Pro Val Val Gln Leu His Ser Asn>	
		1060 1070 1080 1090 1100	
		TTC ACT GCA GTT TGT GTG CTA AAG GAA AAA TGT ATG GAT TAT TTT CAT Phe Thr Ala Val Cys Val Leu Lys Glu Lys Cys Met Asp Tyr Phe His>	
		1110 1120 1130 1140 1150	
		GTA AAT GCT AAT TAC ATT GTC TGG AAA ACA AAC CAT TTT ACT ATT CCT Val Asn Ala Asn Tyr Ile Val Trp Lys Thr Asn His Phe Thr Ile Pro>	
	•	1160 1170 1180 1190 1200	
			٠

Figure 25C

AAG GAG CAA TAT ACT ATC ATA AAC AGA ACA GCA TCC AGT GTC ACC TTT Lys Glu Gln Tyr Thr Ile Ile Asn Arg Thr Ala Ser Ser Val Thr Phe>
1210 1220 1230 1240
* * * * * * * * * * * * * * * * * * *
1250 1260 1270 1280 1290
* * * * * * * * * * TAT GGA ATC ACA ATA ATT TCA GGC TTC GGA CAG CAT GAA CAG AAT GTT TAT GGA ATC ACA ATA ATT TCA GGC Phe Gly Gln Leu Glu Gln Asn Val Tyr Gly Ile Thr Ile Ile Ser Gly>
1300 1310 1320 1330 1340
* * * * * TTG CCT CCA GAA AAA CCT AAA AAT TTG AGT TGC ATT GTG AAC GAG GGG Leu Pro Pro Glu Lys Pro Lys Asn Leu Ser Cys Ile Val Asn Glu Gly>
1350 1360 1370 1380 1390
* * * * * * * * * * * * * * * * * * *
1400 1410 1420 1430 1440
* * * * * * * * * * * * * * * * * * *
1450 1460 1470 1480
* * * * * TOT TOT TOT TOT TOT AAA GCA AAA CGT GAC ACC CCC ACC TCA TGC ACT GTT GAT TAT TCT CYS Lys Ala Lys Arg Asp Thr Pro Thr Ser Cys Thr Val Asp Tyr Ser>
1490 1500 1510 1520 1530
* * * * * * * * * * * * * * * * * * *
1540 1550 1560 1570 1580
* * * * * * * * * * * * * * * * * * *
1590 1600 1610 1620 1630
* * * * * * * * * * * * * * * * * * *
1640 1650 1660 1670 1680
* * * * * * * * * * * * * * * * * * *
1690 1700 1710 1720
* * * * * * * * * * * * * * * * * * *
1730 1740 1750 1760 1770
* * * * * * * * * * ACT TGG AGC CAG ATT CCT CCT GAA GAC ACA GCA TCC ACC CGA TCT TCA ACT TCP Ser Gln Ile Pro Pro Glu Asp Thr Ala Ser Thr Arg Ser Ser

Figure 25D

				•	
1780	1790	1800	18	10	1820
* - TTC ACT (* * * GTC CAA GAC (* * * CTT AAA CCT	TTT ACA GAA	TAT GTG T	TT AGG ATT
Phe Thr	Val Gln Asp	Leu Lys Pro	Phe Thr Glu	Tyr Val P	he Arg Ile>
1830	184) 18	350	1860	1870
* *	* ATG AAG GAA	* * CAT GGT AAG	GGA TAC TGC	AGT GAC	rgg agt gaa
Arg Cys	Met Lys Glu	Asp Gly Lys	Gly Tyr Tri	Ser Asp	Trp Ser Glu>
18	380 . 1	890	1900	1910	1920 .
*	* * AGT GGG ATC	* * * ACC TAT .GAA	* GAT AGA CC	* TCT AAA	GCA CCA AGT
Glu Ala	Ser Gly Ile	Thr Tyr Glu	Asp Arg Pr	o Ser Lys	Ala Pro Ser>
	1930	1940	1950	196	0
*	* * * TAT AAA ATA	* GAT CCA TCC	* * * CAT ACT CA	A GGC TAC	AGA ACT GTA
Phe Trp	Tyr Lys Ile	Asp Pro Ser	His Thr Gl	n Gly Tyr	Arg Thr Val>
1970	1980	1990	2000) 2	2010
<u>.</u>	* * GTG TGG AAG	* * *	* የ ጉርጥ ጥገግ ብ	* AA GCC AAT	GGA AAA ATC
Gln Lev	; GIG IGG AAG 1 Val Trp Lys	Thr Leu Pro	o Pro Phe G	lu Ala Asn	Gly Lys Ile>
2020	2030	. 204		2050	2060
4	* * TAT GAA GTO	* ጉ አርጥ ርጥር ሕር	*	* * AA TCA CAT	TTA CAA AAT
TTG GA: Leu Asj	r TAT GAA GIO p Tyr Glu Va!	Thr Leu Th	r Arg Trp L	ys Ser His	Leu Gln Asn>
207		•	2090	2100	2110
*	* *	* * * * * * * * * * * * * * * * * * * *	* * * * * * * * * * * * * * * * * * *	AT CTC ACA	A AAT GAT CGC
TAC AC Tyr Th	r Val Asn Al	a Thr Lys Le	eu Thr Val A	asn Leu Thi	Asn Asp Arg>
	2120	2130	2140	2150	2160 * *
, * mam_cq	N	 አ አሮኔ ርጥል ል!	*	* * * GTT GGC AA	A TCA GAT GCA
Tyr Le	eu Ala Thr Le	u Thr Val A	rg Asn Leu	Val Gly Ly	s Ser Asp Ala>
	2170	2180	2190	*	200
	* * * ጥጥ ጥጥል አርጥ Aባ	*	* * GT GAC TTT	CAA GCT AC	T CAC CCT GTA
Ala V	al Leu Thr I	le Pro Ala C	ys Asp Phe	Gln Ala Th	r His Pro Val>
2210	2220	2230	22		2250
* ATG G	* * AT CTT AAA G	* CA TTC CCC !	AAA GAT AAC	ATG CTT TO	GG GTG GAA TGG
Met A	sp Leu Lys A	la Phe Pro I	Lys Asp Asn	Met Leu T	rp Val Glu Trp>
2260		0 2:	280	2290	2300 * · *
ACT A	* * ACT CCA AGG G	አኔ ጥርጥ ርጥል	AAG AAA TAT	ATA CTT G	AG TGG TGT GTG
Thr	Thr Pro Arg (Slu Ser Val	Lys Lys Tyr	lle Leu G	
	310	2320	2330	2340 *	2350 * *
* TTA	TCA GAT AAA	GCA CCC TGT	ATC ACA GAC	TGG CAA	CAA GAA GAT GGT
Leu	Ser Asp Lys	Ala Pro Cys			Sin Glu Asp Gly
	2360	2370	2380	239	90 2400

Figure 25

* * * * * * * * * * * * * * * * * * *
2410 2420 2430 2440
* * * * * * * TAT TTG ATA ACA GTT ACT CCA GTA TAT GCT GAT GGA CCA GGA AGC CCT Tyr Leu Ile Thr Val Thr Pro Val Tyr Ala Asp Gly Pro Gly Ser Pro>
2450 2460 2470 2480 2490
GAA TCC ATA AAG GCA TAC CTT AAA CAA GCT CCA CCT TCC AAA GGA CCT Glu Ser Ile Lys Ala Tyr Leu Lys Gln Ala Pro Pro Ser Lys Gly Pro>
2500 2510 2520 2530 2540
ACT GTT CGG ACA AAA AAA GTA GGG AAA AAC GAA GCT GTC TTA GAG TGG Thr Val Arg Thr Lys Lys Val Gly Lys Asn Glu Ala Val Leu Glu Trp>
2550 2560 2570 2580 2590 * * * * * * * * *
GAC CAA CTT CCT GTT GAT GTT CAG AAT GGA TTT ATC AGA AAT TAT ACT Asp Gln Leu Pro Val Asp Val Gln Asn Gly Phe Ile Arg Asn Tyr Thr>
2600 2610 2620 2630 2640
ATA TTT TAT AGA ACC ATC ATT GGA AAT GAA ACT GCT GTG AAT GTG GAT Ile Phe Tyr Arg Thr Ile Ile Gly Asn Glu Thr Ala Val Asn Val Asp>
2650 2660 . 2670 2680
TCT TCC CAC ACA GAA TAT ACA TTG TCC TCT TTG ACT AGT GAC ACA TTG Ser Ser His Thr Glu Tyr Thr Leu Ser Ser Leu Thr Ser Asp Thr Leu>
2690 2700 2710 2720 2730
TAC ATG GTA CGA ATG GCA GCA TAC ACA GAT GAA GGT GGG AAG GAT GGT Tyr Met Val Arg Met Ala Ala Tyr Thr Asp Glu Gly Gly Lys Asp Gly>
2740 2750 2760 2770 2780
CCA GAA TTC ACT TTT ACT ACC CCA AAG TTT GCT CAA GGA GAA ATT GAA Pro Glu Phe Thr Phe Thr Pro Lys Phe Ala Gln Gly Glu Ile Glu>
2790 2800 2810 2820 2830
* * * TOO GGG GGC GAC AAA ACT CAC ACA TGC CCA CCG TGC CCA GCA CCT GAA Ser Gly Gly Asp Lys Thr His Thr Cys Pro Pro Cys Pro Ala Pro Glu>
2840 2850 2860 2870 2880
CTC CTG GGG GGA CCG TCA GTC TTC CTC TTC CCC CCA AAA CCC AAG GAC Leu Leu Gly Gly Pro Ser Val Phe Leu Phe Pro Pro Lys Pro Lys Asp>
2890 2900 2910 2920
ACC CTC ATG ATC TCC CGG ACC CCT GAG GTC ACA TGC GTG GTG GAC Thr Leu Met Ile Ser Arg Thr Pro Glu Val Thr Cys Val Val Val Asp
2930 2940 2950 2960 2970
GTG AGC CAC GAA GAC CCT GAG GTC AAG TTC AAC TGG TAC GTG GAC GGC

	Val Ser His Glu Asp Pro Glu Val Lys Phe Asn Trp Tyr Val Asp Gly>
	2980 2990 3000 3010 3020
· · · · · · · · · · · · · · · · · · ·	GTG GAG GTG CAT AAT GCC AAG ACA AAG CCG CGG GAG GAG CAG TAC AAC Val Glu Val His Asn Ala Lys Thr Lys Pro Arg Glu Glu Gln Tyr Asn>
· .	3030 3040 3050 3060 3070
•	AGC ACG TAC CGT GTG GTC AGC GTC CTC ACC GTC CTG CAC CAG GAC TGG Ser Thr Tyr Arg Val Val Ser Val Leu Thr Val Leu His Gln Asp Trp>
•	3080 3090 3100 3110 3120
	CTG AAT GGC AAG GAG TAC AAG TGC AAG GTC TCC AAC AAA GCC CTC CCA Leu Asn Gly Lys Glu Tyr Lys Cys Lys Val Ser Asn Lys Ala Leu Pro>
	3130 3140 3150 3160
	GCC CCC ATC GAG AAA ACC ATC TCC AAA GCC AAA GGG CAG CCC CGA GAA Ala Pro Ile Glu Lys Thr Ile Ser Lys Ala Lys Gly Gln Pro Arg Glu>
	3170 3180 3190 3200 3210
	CCA CAG GTG TAC ACC CTG CCC CCA TCC CGG GAT GAG CTG ACC AAG AAC Pro Gln Val Tyr Thr Leu Pro Pro Ser Arg Asp Glu Leu Thr Lys Asn>
	3220 3230 3240 3250 3260
	CAG GTC AGC CTG ACC TGC CTG GTC AAA GGC TTC TAT CCC AGC GAC ATC Gln Val Ser Leu Thr Cys Leu Val Lys Gly Phe Tyr Pro Ser Asp Ile>
	3270 3280 3290 3300 3310
	GCC GTG GAG TGG GAG AGC AAT GGG CAG CCG GAG AAC AAC TAC AAG ACC Ala Val Glu Trp Glu Ser Asn Gly Gln Pro Glu Asn Asn Tyr Lys Thr>
	3320 3330 3340 3350 3360
	ACG CCT CCC GTG CTG GAC TCC GAC GGC TCC TTC TTC CTC TAC AGC AAG Thr Pro Pro Val Leu Asp Ser Asp Gly Ser Phe Phe Leu Tyr Ser Lys>
· .	3370 3380 3390 3400
	CTC ACC GTG GAC AAG AGC AGG TGG CAG CAG GGG AAC GTC TTC TCA TGC Leu Thr Val Asp Lys Ser Arg Trp Gln Gln Gly Asn Val Phe Ser Cys>
	3410 3420 3430 3440 3450
	TCC GTG ATG CAT GAG GCT CTG CAC AAC CAC TAC ACG CAG AAG AGC CTC Ser Val Met His Glu Ala Leu His Asn His Tyr Thr Gln Lys Ser Leu>
·	3460 3470 * *
	TCC CTG TCT CCG GGT AAA TGA Ser Leu Ser Pro Gly Lys ***>
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•	
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		1	.0	٠		20		4.	30		•	4	40	*		
ATG Met	* GTG Val	CTT Leu	* CTG Leu	* TGG Trp	TGT Cys	* GTA Val	GTG Val	AGT Ser	CTC Leu	TAC Tyr	TTT Phe	тат Туг	GGA Gly	ATC Ile	CTG Leu	>
50 * CAA	AGT	* GAT	60 * GCC Ala	TCA	* GAA	CGC	70 * TGC	* GAT	GAC Asp	80 * TGĢ	GGA Gly	* CTA	90 * GAC Asp	ACC	* ATG Met	; .>
		Asp		110	G1u	ALG	120	•			.30		_	140		
	00 * CAA Gln	ATC	CAA Gln	*	TTT Phe	* GAA Glu	* raė	GAG	* CCA Pro	A GCT	* CGC Arg	* OTA :	. AAC	* G TGC	CC?	\ >>
*	150) :	* A CAC a His	1 • • • • • • • • • • • • • • • • • • •	.60 *	* 2 A A .:	ነ	170 *	C TA	* C AG	180 C AC	0 * A GC	* C CA'	r TC	190 * A GC'	${f T}$
Lev	ı Phe		ι Hls	s Pne	21(5 Fil		220			230			24	
GG(* C CT y Le	200 * T AC u Th	T CTO	* G AT	C TC(* ጌ ጥል ⁽	* T TG r Tr	G AC	* T AG	G CA	* .G GA .n As	t C CG p Ar	G GA	* C CT p Le	T GA u Gl	* .G .u>
			250			260		*	27	70		ŧ	280		*	
GA Gl	G CC		* T AA e As	c mu	* C CG e Ar	C CI	C CC	C GA	AG AZ	AC CO	GC AT	TT AC	er Ly	AG G! ys G:	AG AI Lu Ly	AA ys>
290				00		,	310		*	32	0 *	*		30 *		*
GA As	T G T Sp V	rG Ci al Le	rg To eu Ti		nc cc	c cc	CC A	CT C' hr L	TC C eu L	TC A eu A	AT G	AC A sp T	CT G hr G	GC A ly A	AC T sn T	AT>
	340			35	0	*		60 *		*	370 *		*	38	0	
A T	CC T hr C	~~ `	TG T	TA A eu A	GG A rg A	AC A	ርጥ Δ	CA T	'AT T	Cys S	GC A Ser I	JAA (TT G	CA T	rrr c Phe F	ccc ?ro>
		90			400		*	41	LO *		<i>k</i>	120 *		*	430) *
	TG C	* SAA C Slu \	GTT C	* STT (Val (CAA A	AA C ys A	מבי	AGC (Ser (rgr ' Cys	TTC Phe	AAT ' Asn '	TCC (Ser	Pro 1	ATG A Met	AAA Lys	CTC Leu>
		4	10			150		*	46	0 *	*	4	70 *		\ ★	480
(cCA (Pro	GTG (Val	* CAT A	AAA (T CTG ' Leu '	rat / Tyr	ATA Ile	GAA Glu	TAT Tyr	GGC Gly	ATT Ile	CAG Gln	AGG Arg	ATC Ile	ACT Thr	TGT Cys>
•			49	0		5	0.0		*	510		*	52	20 *	*	
	CCA Pro	* AAT Asn	GTA Val	* GAT Asp	GGA Gly	TAT ·	TTT Phe	CCT Pro	TCC Ser	AGT Ser	GTC Val	AAA Lys	CCG Pro	ACT Thr	ATC Ile	ACT Thr>
5	30		*	540		*	5!	50 *	*	ļ	560	-	*.	570 *		*
	TGG	TAT	ATG	GGC	TGT	TAT	AAA	ATA Ile	CAG Gln	AAT Asn	TTT Phe	AAT Asn	AAT Asn	GTA Val	ATA Ile	CCC Pro

Figure 26B

•	580 590	600	610. * *	620 * *	
	GAA GGT ATG AAC TTG AGIu Gly Met Asn Leu S	GT TTC CTC ATT Ser Phe Leu Ile	GCC TTA ATT TC Ala Leu Ile Se	A AAT AAT GGA r Asn Asn Gly>	
	630 640	650	660	670 * *	
	AAT TAC ACA TGT GTT (Asn Tyr Thr Cys Val	GTT ACA TAT CCA Val Thr Tyr Pro	GAA AAT GGA CO	T ACG TTT CAT Tg Thr Phe His>	
			00 710		
	* * * CTC ACC AGG ACT CTG Leu Thr Arg Thr Leu	* * ACT GTA AAG GTA Thr Val Lys Val	* * * * * * * * * * * * * * * * * * *	CA AAA AAT GCA ro Lys Asn Ala>	
	730	740	750	760 .	
	* * * * GTG CCC CCT GTG ATC Val Pro Pro Val Ile	CAT TCA CCT AAT His Ser Pro Asi	r GAT CAT GTG G n Asp His Val V	TC TAT GAG AAA al Tyr Glu Lys>	
•	770 780	790	* * *	810	•
	GAA CCA GGA GAG GAG Glu Pro Gly Glu Glu	CTA CTC ATT CC	C TGT ACG GTC To Cys Thr Val T	PAT TTT AGT TTT Tyr Phe Ser Phe>	•
•	820 830	840 * *	850 * *	860 * *	
	CTG ATG GAT TCT CGC Leu Met Asp Ser Arg	AAT GAG GTT TO Asn Glu Val Tr	GG TGG ACC ATT of Trp Trp Thr Ile	GAT GGA AAA AAA Asp Gly Lys Lys>	,
	* *	80 890 * *	k * *	910 * .*	
	CCT GAT GAC ATC ACT	ATT GAT GTC AC	cc ATT AAC GAAhr Ile Asn Glu	AGT ATA AGT CAT Ser Ile Ser His>	
	920	930	* *	* * * *	
•	AGT AGA ACA GAA GA' Ser Arg Thr Glu As	r GAA ACA AGA A p Glu Thr Arg T	CT CAG ATT TTG	AGC ATC AAG AAA Ser Ile Lys Lys>	•
	970	980 * * *	990	1000	
	GTT ACC TCT GAG GA Val Thr Ser Glu As	T CTC AAG CGC A p Leu Lys Arg S	GC TAT GTC TGT Ser Tyr Val Cys	CAT GCT AGA AGT His Ala Arg Ser>	
	1010 1020	1030 * *	1040	1050 * * *	•
	GCC AAA GGC GAA GT Ala Lys Gly Glu Va	TT GCC AAA GCA (al Ala Lys Ala i	GCC AAG GTG AAG Ala Lys Val Lys	CAG AAA GTG CCA Gln Lys Val Pro>	
	1060 * 1070	* * *	1090 * *	1100 * *	
	GCT CCA AGA TAC AGA Ala Pro Arg Tyr T	CA GTG TCC GGT hr Val Ser Gly	GGC GCG CCT ATO	CTG AGC GAG GCT Leu Ser Glu Ala>	
	* * *	* *	30 * 114	* * *	
	GAT AAA TGC AAG G Asp Lys Cys Lys G	AA CGT GAA GAA lu Arg Glu Glu	AAA ATA ATT TT. Lys Ile Ile Le	A GTG TCA TCT GCA u Val Ser Ser Ala>	
	1160	1170	1180	1190	•
			·		
	•		•		
<i>:</i>					
	•	•			

AAT GAA ATT GAT GTT CGT CCC TGT CCT CTT AAC CCA AAT GAA CAC AAA

AAT GAA ATT GAT GIT COT COS Pro Leu Asn Pro Asn Glu His Lys> Asn Glu Ile Asp Val Arg Pro Cys Pro Leu Asn Pro Asn Glu His Lys>
1210 1220 1230 1240
* * * * * * * * * * * * * * * * * * *
1250 1260 1270 1280 1290
* * * * * * * * * * * * * * * * * * *
1300 1310 1320 1330 1340
* * * * * * * CCT GCT AAG GTG GAG GAT TCA GGA CAT TAC TAT TGC GTG GTA AGA AAT CCT GCT AAG GTG GAG GAT TCA GGA CAT TAC TAT TGC GTG GTA AGA AAT CCT GCT GAG GTG GTA AGA AAT Pro Ala Lys Val Glu Asp Ser Gly His Tyr Tyr Cys Val Val Arg Asn>
1350 1360 1370 1380 1390
* * * * * * * * * * * * * * * * * * *
1400 1410 1420 1430 1440
* * * * * * * * * * GCA CAA GCC ATA TTT AAG CAG AAA CTA GLU Pro Asn Leu Cys Tyr Asn Ala Gln Ala Ile Phe Lys Gln Lys Leu>
1450 1460 1470 1480
* * * * * * * * * * * * * * * * * * *
1490 1500 1510 1520 1530
* * * * * * * * * * * * * * * * * * *
1540 1550 1560 1570 1580 * * * * * * * * * * * * * * * * * * *
* * * * * * * * * * * * * * * * * * *
1590 1600 1610 1620 1630 * * * *
* * * * * * * * * * * * * * * * * * *
1640 1650 1660 1670 1680
* * * * * * * * CAT GCA TAC TAC TTG GGC AAG CAA TAT CCT ATT ACC CGG GTA CAT GCA TCC TAC ACA TAC TTG GGC AAG CAA TAT CCT ATT ACC CGG GTA His Ala Ser Tyr Thr Tyr Leu Gly Lys Gln Tyr Pro Ile Thr Arg Val:
1690 1700 1710 1720
ATA GAA TTT ATT ACT CTA GAG GAA AAC AAA CCC ACA AGG CCT GTG ATT Ile Glu Phe Ile Thr Leu Glu Glu Asn Lys Pro Thr Arg Pro Val Ile
1730 1740 1750 1760 1770
* * * * * * GTG AGC CCA GCT AAT GAG ACA ATG GAA GTA GAC TTG GGA TCC CAG ATA

Figure 26D

	1780 1790 1800 1810 1	820
	CAA TTG ATC TGT AAT GTC ACC GGC CAG TTG AGT GAC ATT GCT Gln Leu Ile Cys Asn Val Thr Gly Gln Leu Ser Asp Ile Ala	TAC TGG Tyr Trp>
•	1830 1840 1850 1860	1870
	AAG TGG AAT GGG TCA GTA ATT GAT GAA GAT GAC CCA GTG CTA Lys Trp Asn Gly Ser Val Ile Asp Glu Asp Asp Pro Val Leu	GGG GAA Gly Glu>
	1880 1890 1900 1910	1920
	GAC TAT TAC AGT GTG GAA AAT CCT GCA AAC AAA AGA AGG AGG AGG ASP Tyr Tyr Ser Val Glu Asn Pro Ala Asn Lys Arg Arg Ser	r ACC CTC r Thr Leu>
	1930 1940 1950 1960 * * * * * * * * *	. *
	ATC ACA GTG CTT AAT ATA TCG GAA ATT GAG AGT AGA TTT TA Ile Thr Val Leu Asn Ile Ser Glu Ile Glu Ser Arg Phe Ty	T AAA CAT r Lys His>
•	1970 1980 1990 2000 201	0 * *
	CCA TTT ACC TGT TTT GCC AAG AAT ACA CAT GGT ATA GAT GC Pro Phe Thr Cys Phe Ala Lys Asn Thr His Gly Ile Asp Al	A GCA TAT a Ala Tyr>
	2020 2030 2040 2050	2060
	ATC CAG TTA ATA TAT CCA GTC ACT AAT TCC GGA GAC AAA AG Ile Gln Leu Ile Tyr Pro Val Thr Asn Ser Gly Asp Lys T	or His Thr>
•	2070 2080 2090 2100 * * * * * * * *	2110 * *
	TGC CCA CCG TGC CCA GCA CCT GAA CTC CTG GGG GGA CCG T Cys Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro S	CA GTC TTC er Val Phe>
	2120 2130 2140 2150	* *
	CTC TTC CCC CCA AAA CCC AAG GAC ACC CTC ATG ATC TCC C Leu Phe Pro Pro Lys Pro Lys Asp Thr Leu Met Ile Ser A	GG ACC CCT Arg Thr Pro>
•	2170 2180 2190 ' 2200 * * * * * * * * *) * *
	GAG GTC ACA TGC GTG GTG GTG GAC GTG AGC CAC GAA GAC Glu Val Thr Cys Val Val Val Asp Val Ser His Glu Asp	CCT GAG GTC Pro Glu Val>
·	2210	250 * *
•	AAG TTC AAC TGG TAC GTG GAC GGC GTG GAG GTG CAT AAT Lys Phe Asn Trp Tyr Val Asp Gly Val Glu Val His Asn	GCC AAG ACA Ala Lys Thr>
	2260 2270 2280 2290	2300
	AAG CCG CGG GAG GAG CAG TAC AAC AGC ACG TAC CGT GTG Lys Pro Arg Glu Glu Gln Tyr Asn Ser Thr Tyr Arg Val	GTC AGC GTC Val Ser Val>
•	2310 2320 2330 2340	2350 * *
	CTC ACC GTC CTG CAC CAG GAC TGG CTG AAT GGC AAG GAG Leu Thr Val Leu His Gln Asp Trp Leu Asn Gly Lys Glu	TAC AAG TGC Tyr Lys Cys>
•	2360 2370 2380 2390	2400
· · · .		

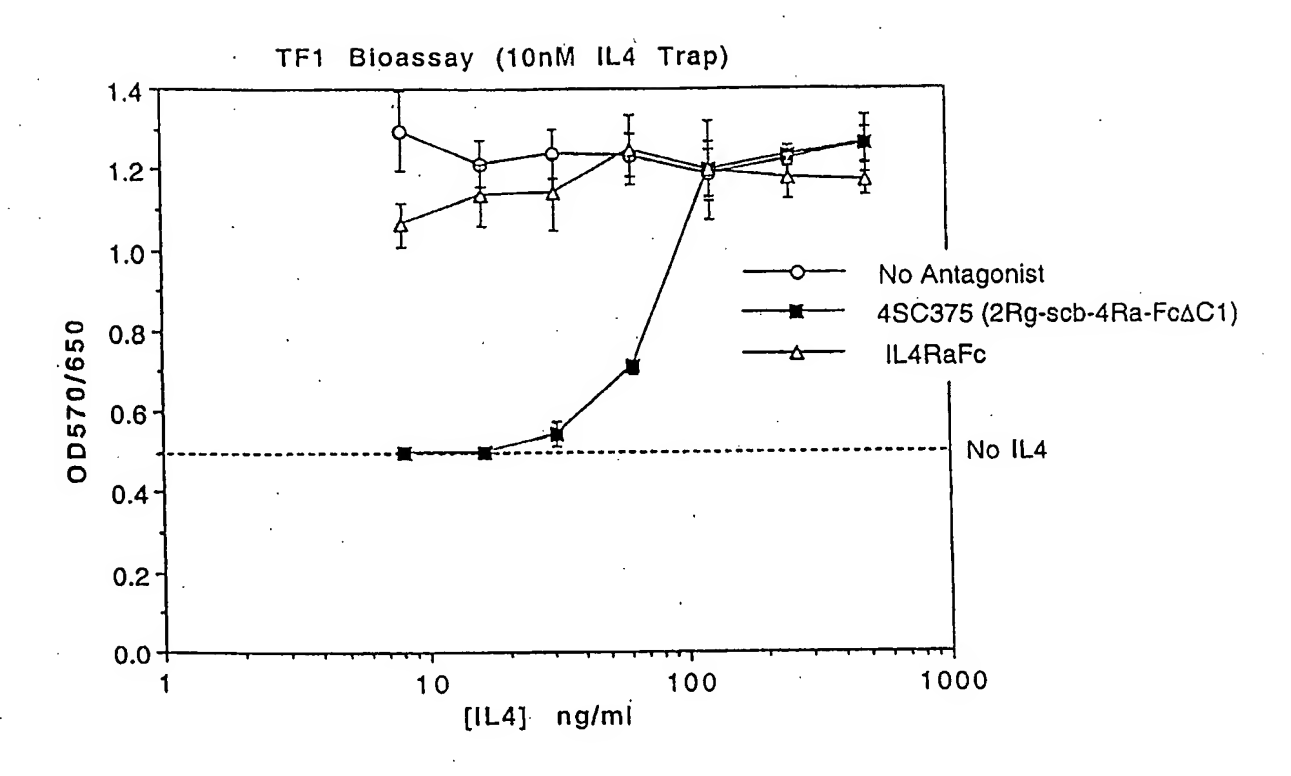
51/74

Figure 26E

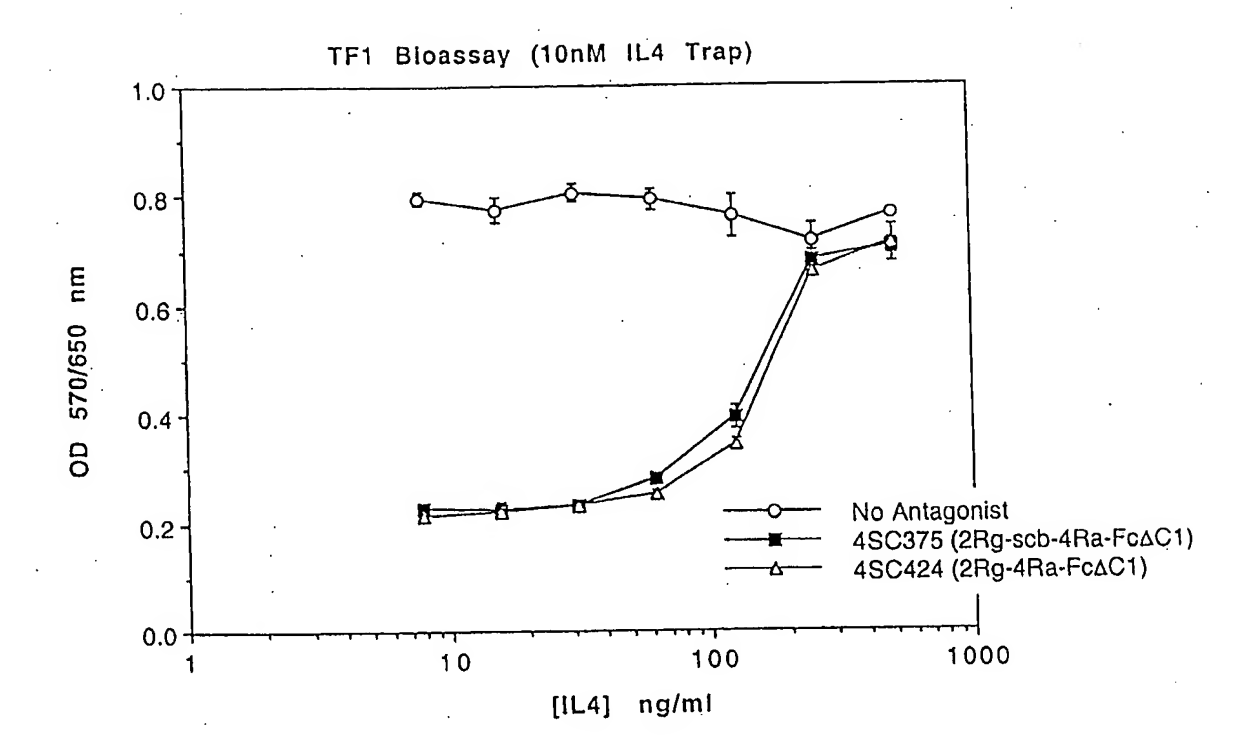
	*		*		*	*		*		*	*		*		*	*
	N N C	CMC	mCC	אאר	מממ	GCC	ርጥር	CCA (GCC	CCC	ATC	GAG	AAA	ACC	ATC	TCC
	AAG	Ual I	202	AAC	ING	Ala	Leu	Pro	Ala	Pro	Ile	Glu	Lys	Thr	Ile	Ser>
	nys .	VQI	DET	No!!	273	1120	200									
			24	10		2	120		2	430			244	0		
		*		*	*		*		*	*		*		*	*	
	AAA	GCC	AAA	GGG	CAG	CCC	CGA	GAA	CCA	CAG	GTG	TAC	ACC	CTG	CCC	CCA
	Lys	Ala	Lys	Gly	Gln	Pro	Arg	Glu	Pro	Gln	Val	Tyr	Thr	Leu	Pro	Pro>
										•	•					
24	150			2460	ı		24	70		2	480		<u>.</u>	2490		*
	*		*	*		*	•	*	*		*	OMO.	x 200	: " maa	OMO	
	TCC	CGG	GAC	GAC	ATG	ACC	· AAG	AAC	CAG	GTC	AGC	CTG	ACC	TGC	CTG	GIC .
	Ser	Arg	Glu	Glu	Met	Thr	ГЛЗ	Asn	Gin	val	Ser	Leu	THE	Cys	neu	Val>
		a.a			\ .			2520			25	3.0		2	540	
	25	00		L	2510		*	2320 *		*		*	*	-	*	•
		*		י יי מיאו	n aca	ነ አሮር	. СУС	ልጥ ር	GCC	GTG	GAG	TGG	GAG	AGC	AAT	GGG
	AAA	GG(TIT	J TA	r Dro	. AGC	. Asn	Tle	Ala	Val	Glu	Trp	Glu	Ser	Asn	Gly>
	rys	GTA	PIN	: = 1	r Pr(, ind	#10				_				
		2550	1		25	560		2	570	•		2580	•		25	90
	•		k	*		*	*		*		*	*		*		*
	CAG	CCC	G GA	g aa	C AA	C TAC	DAA C	ACC	ACG	CCI	ccc	GTG	CTG	GAC	TCC	GAC
	Gln	Pro	o G1	u As	n Asi	n Ty	r Lys	Thr	Thr	Pro) Pro	Val	. Leu	Asp	Ser	<qaa< td=""></qaa<>
						•										
	•		2600			261	0		26	520			2630		· *	2640
	4	•	*		*	. 4	*	*	· cm	*;						
	GGC	TC	C TI	C TI	C CT	C TA	T AGO	AAG	CYC	. Mh	v Va	G GAG	- AAK	s AGC	r Arc	G TGG
	Gly	/ Se	r Ph	e Pr	e Le	u Ty	r Se	с гуз	; Le	7 771	ı va	T WO	o my a	,		g Trp>
			,) C E O			2660			267	0		20	580		
		•		8650		*	*		*		*	*		*		*
	C N (״ גיט בי	c co	מב או	የር G1	יר יד	C TC	A TG	TC	C GT	G AT	G CA	T GA	G GC'	T CT	G CAC
	GI	n Gl	n G	V A	sn Va	al Ph	e Se	r Cys	s Se	r Va	l Me	t Hi	s Gl	u Al	a Le	u His>
	U			-,,		_		-		,					_	
	2690			27	00		2	710			2720		_	273	0	
	4		*		*	1		*		*	†		*	m = =	* **	13
	AA	C CA	C T	AC A	CG C	AG AA	G AG	C CT	C TC	C C1	rg To	CT CC	G GG	'I' AA	A TO	6A : * <
	Ac	n Hi	is m	vr T	hr G	ln Ly	ys Se	r Le	u Se	er Le	eu Se	er Pr	O GT	У гу	່ ສຳ	~ >

PCT/US99/22045

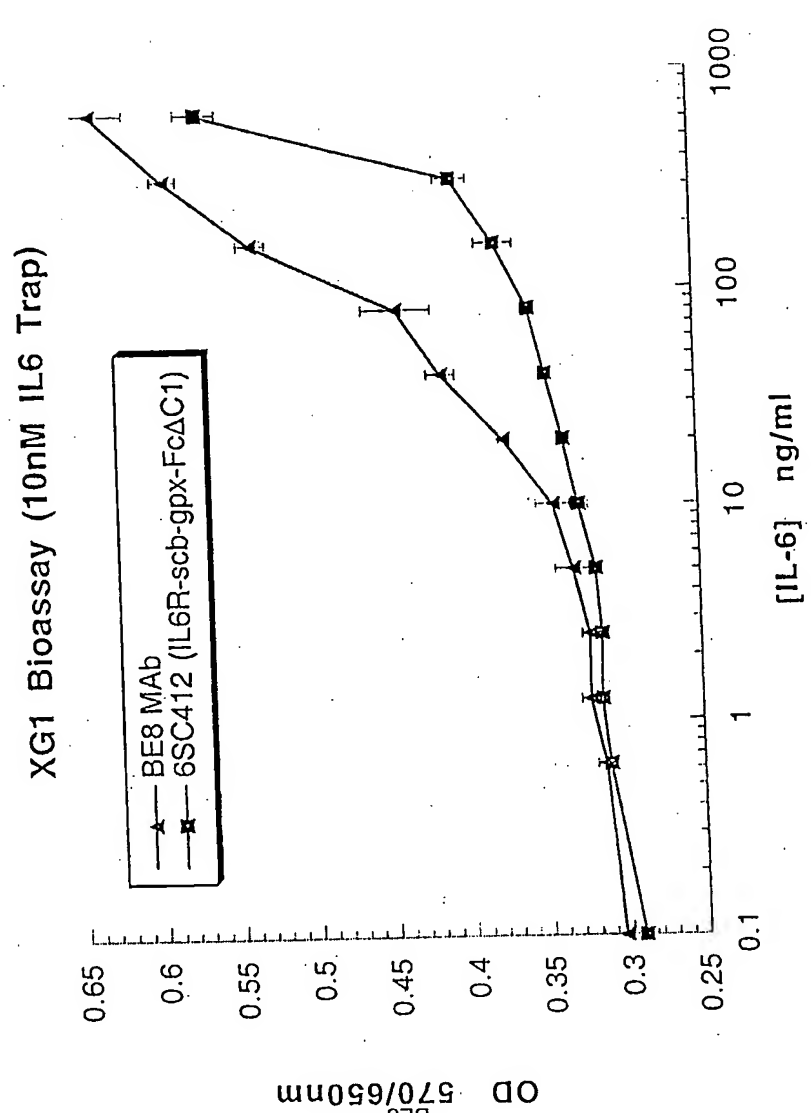
52/74 Figure 27



53/ 74 Figure 28



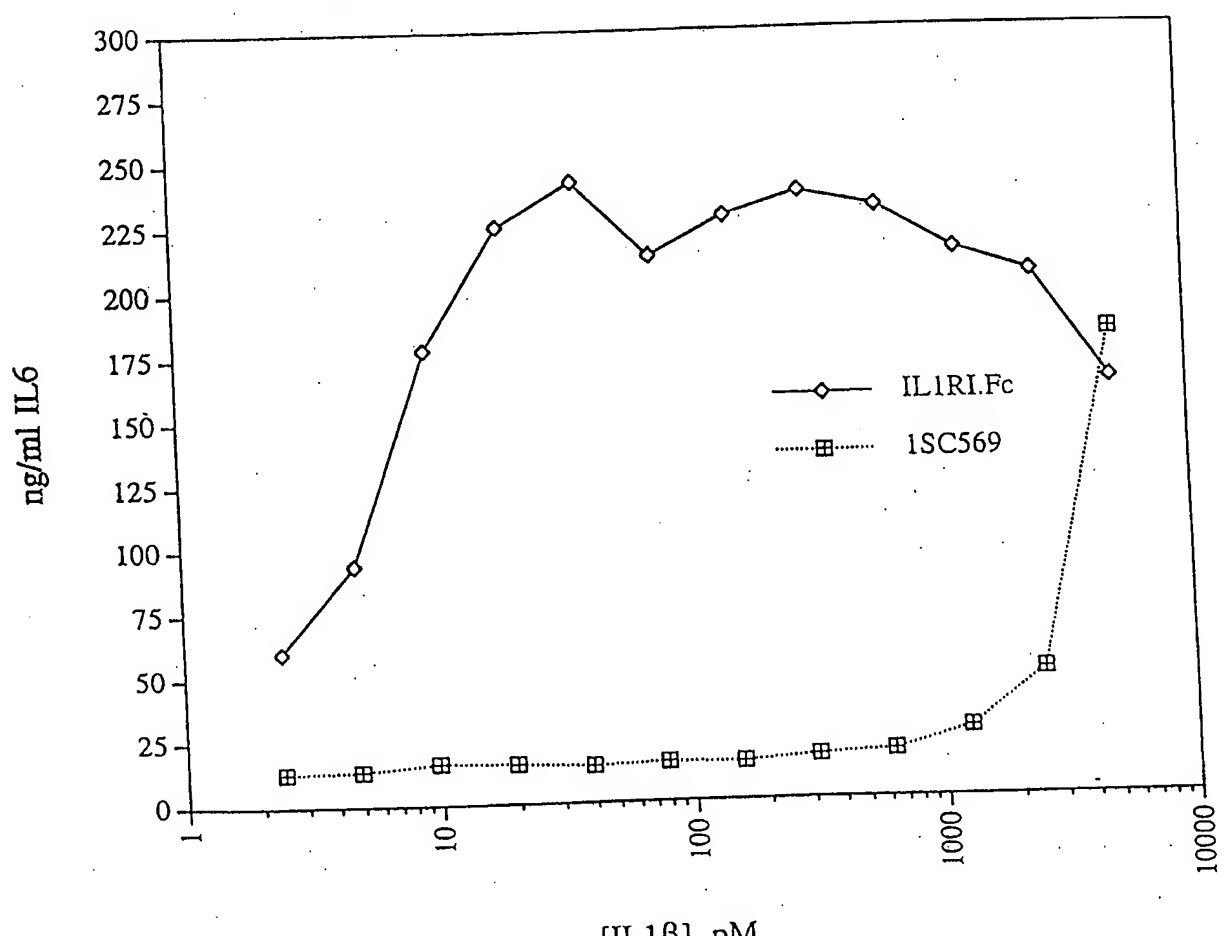
54/74 Figure 29



270/650nm BE8

55/ **7**4 Figure 30

MRC5 Bioassay (10nM IL1 Trap)
IL1 Trap 1SC569 vs IL1 Trap IL1RI.Fc



[IL1 β] pM

Figure 31A

10		20 .			40
	*				* * .
ATG GTG TGG C	TT TGC TCT	GGG CTC	CTG TTC	CCT GTG A	GC TGC CTG GTC
TAC CAC ACC G	AA ACG AGA	CCC GAG	GAC. AAG	Pro Val S	CG ACG GAC CAG er Cys Leu Val>
Met Val Trp L	seu Cys Ser	GIA men	Leu File	PLO VAI S	cr cyb bed var
50	60	70		80	90
* *	* *				* *
CTG CTG CAG	TG GCA AGC	TCT GGG	AAC ATG	AAG GTC T	TG CAG GAG CCC
GAC GAC GTC C	CAC CGT TCG	AGA CCC	TTG TAC	TTC CAG A	AC GTC CTC GGG
Leu Leu Gln V	/al Ala Ser	Ser Gly	Ash Met	nys var n	eu Gln Glu Pro>
100	110	120		130	140.
* . *	*	* *	*	*	* *
ACC TGC GTC	rcc gac tac	ATG AGC	ATC TCT	ACT TGC G	AG TGG AAG ATG
TGG ACG CAG	AGG CTG ATG	TAC TCG	TAG AGA	TGA ACG	TC ACC TTC TAC
Thr Cys Val	Ser Asp Tyr	Met Ser	Ile Ser	Thr Cys G	Slu Trp Lys Met>
150	160		170	180	190
	* *	*	*	* *	* *
AAT GGT CCC	ACC AAT TGO	AGC ACC	GAG CTC	CGC CTG	TTG TAC CAG CTG
TTA CCA GGG	TGG TTA ACC	TCG TGG	CTC GAG	GCG GAC	AAC ATG GTC GAC
Asn Gly Pro	Thr Asn Cys	s Ser Thr	Glu Leu	Arg Leu I	Leu Tyr Gln Leu>
200	210	1	220	2:	30 240
* *					* * *
GTT TTT CTG	CTC TCC GA	A GCC CAC	ACG TGT	ATC CCT	GAG AAC AAC GGA
CAA AAA GAC	GAG AGG CT	r CGG GTG	TGC ACA	TAG GGA	CTC TTG TTG CCT
Val Phe Leu	Leu Ser Gl	u Ala His	Thr Cys	s Ile Pro	Glu Asn Asn Gly>
25	50	260	. 270)	280
*					* *
GGC GCG GGG	TGC GTG TG	C CAC CTO	G CTC ATC	G GAT GAC	GTG GTC AGT GCG
CCG CGC CCC	ACG CAC AC	G GTG GAG	C GAG TAG	C CTA CTG	CAC CAG TCA CGC
Gly Ala Gly	Cys Val Cy	s His Le	u Leu Met	t Asp Asp	Val Val Ser Ala>
0.00	300	310.		320	330
290 .	* *	*	*	*	* * *
GAT AAC TAT	ACA CTG GA	C CTG TG	G GCT GG	G CAG CAG	CTG CTG TGG AAG
CTA TTG ATA	TGT GAC CI	G GAC AC	C CGA CC	C GTC GTC	GAC GAC ACC TTC
Asp Asn Tyr	Thr Leu As	sp Leu Tr	p Ala Gl	y Gln Gln	Leu Leu Trp Lys>
2.4.0	350	36	0	370	380
340	350 *	*	* *		* *
GGC TCC TTC	AAG CCC AG	GC GAG CA	T GTG AA	A CCC AGG	GCC CCA GGA AAC
CCG AGG AAG	TTC GGG TO	CG CTC GI	A CAC TI	T GGG TCC	CGG GGT CCT TTG
Gly Ser Phe	Lys Pro S	er Glu Hi	s Val Ly	s Pro Arg	Ala Pro Gly Asn>

Figure 31B

		390			400	0	4.	4	10		.	420		•	430)			
	* CTG /	* >~	<u>~~~~ /</u>	· ች ጉአጠ ·	አሮሮ	አአጥ ·	- አ መር	TCC .	CAC	ልሮጥ	° СФС	CTC.	ርጥር	` እርር <i>'</i>	TGG 1	7.GC			
	GAC '																		
	Leu '																		
	Leu	TIIT	vai i	1113	1114 4	Non	Var				<u>س</u> وت							•	
		4	40		,	450			46	0		4	70			480		•	
	*		*		.*	*		*		*	*		. *		. *	*			
, ·	AAC	CCG	TAT	CCC	CCT	GAC	AAT	TAC	CTG	TAT	TAA	CAT	CTC	ACC	TAT	GCA			
	TTG																•		
	· Asn	Pro	Tyr	Pro	Pro	Asp	Asn	Tyr	Leu	Tyr	Asn	His	Leu	Thr	Tyr	Ala>			
				•		_				C 1 A			E 2	^					
		*	49	() *	*	ל	\$00 *		★.	510 *		*	. 52	*	*				
·	GTC	AAC	ATT	TGG	AGT	GAA	AAC	GAC	CCG	GÇA	GAT	TTC	AGA	ATC	TAT	AAC	•		
•	CAG																		
	Val																		
				_						_	مدامر بر مدامر بر								
	530			540			. 55	50		į	560		ı.	570			•		
	*	. ~ ~	* ·	*	71.	*		T C C C	*	z ma	~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	CCC	* * カロロ	א חחת	CIDC	^አ			
													AGC TCG						
•																Lys>		•	
	Val	1111	тÄт	nea	GIU	PIO	201	Beu	•••							2			
	. 58	30		. '	590			600			6	10	•	(620		_		
		*	*		*		*	*		*		*	*		*		•		
													GCT						
													CGA						
•	Ser	Gly	Ile	Ser	Tyr	Arg	Ala	Arg	Val	Arg	Ala	Trp	Ala	Gln	Ser	Tyr>			
•		<i></i>			سر	4.0		•	CEO			660	•		۲,	70			
	4	630		*	0	40 *	*		650 *		*	*		*	U	*			
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													ACC				•		
•																Ser>			
•	¥			-															
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													G GGC						
•																GGA Pro>			
	TYE	wig	GIU	, rrc) Elle	S GT(r ATT	. vel	. Grj	. ar7	, O.L.	ı Or.	, U±)						
			7	30	•		740			750)			760					
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																AAC			
,																TTG			
	Thr	Gli	ı Thr	Gli	n Pro		o Val		r Ası	n Le	u Se	r Va	ı Se:	r va.	ı Gil	Asn>			
							•			•		•							
								•			•								
•																			
					•														
		•																	

Figure 31C

770	780					79	0		8	00					
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			GTA												
			CAT												
Leu	Суѕ	Thr	Val	11e	Trp	Thr	Trp	Asn	PIO	PIO	Giu	GīĀ	мта	ser	261 >
82	20		8	30			840			85	0		8	60	
	*	*		*		*	*		*		*	*		*	
		_													AAG ·
			GAT												
Asn	Cys	Ser	Leu	Tip	TAT	Pne	Ser	UTP	rne	Gry	Map	цуз	GTII	vaħ	Dy 37
	870	•		88	80		8	90			900			91	.0
*	*	•				` *		*,	•	*	*		* .		*
			CCG				•								
			GGC												
Lys	Ile	Ala	Pro	Glu	Thr	Arg	Arg	Ser	IIe	GIU	vaı	Pro	ren	Asn	Glu>
	ı	920			930			9.	40		9	950			960
*		.*		*	*		*		*	*		*		*	*
AGG	ATT	TGT	CTG	CAA	GTG	GGG	TCC	CAG	TGT	AGC	ACC	AAT	GAG	AGT	GAG
			GAC												
Arg	Ile	Cys	Leu	Gln	Val	Gļy	Ser	Gln	Cys	Ser	Thr	Asn	Glu	Ser	Glu>
		q	70		•	980			990			10	00		
	*		*	*	•	*		*	*		*			*	
AAG	CCT	AGC	ATT	TTG	GTT	GAA	AAA	TGC	ATC	TCA	CCC	CCA	GAA	GGT	GAT
TTC	GGA	TCG	TAA	AAC	CAA	CTT	TTT	ACG	TAG	AGT	GGG	GGT	CTT	CCA	CTA
Lys	Pro	Ser	Ile	Leu	Val	Glu	Lys	Cys	Ile	Ser	Pro	Pro	Glu	Gly	Asp>
1010			1020			1.0	30		1	040			1050		
*			*												*
CCT	' GAG	TCI	GCT	GTG	ACT	GAG	CTT	CAA	TGC	TTA	TGG	CAC	AAC	CTG	AGC .
			CGA												
Pro	Glu	ı Ser	Ala	Val	Thr	Glu	Leu	Glr	Cys	s Ile	Trp	His	Asn	Leu	Ser>
. 10	160		1	070			1080			10	90		1	.100	
20	*		k											*	
															ACT
															TGA
Туг	: Met	Lys	з Суз	Ser	Trp	Lev	Pro	Gly	y Aro	g Asr	ı Thr	Sei	Pro) Asr	Thr>
	1110	n		11	20		1	.130		•	1140)		11	.50
*		*	* .			4				*	-		*		
AA(TA	r AC'	r cro	TAC	TAT	r ŤG(G CAC	AG	A AG	C CTO	GA/	AA A	TA A	r car	r caa
TT	G AT	A TG	A GAG	TA E	ATA	A AC	GTO	TC'	T TC	G GA	CT	r TT'	TAI	A GT	A GTT
Ası	n Ty	r Th	r Lei	ı Tyı	туз	r Tri	o His	s Ar	g Se	r Le	ı Glı	ı Ly	s Il	e Hi	s Gln>

Figure 31D

	1160	1	170	1180	1190	1200	
•	* *	, *	* *	k *	* *	* *	
	TGT GAA AAC	ATC TTT	AGA GAA GO	GC CAA TAC TT	T GGT TGT TCC	TTT GAT	
	ACA CTT TTG	TAG AAA '	TCT CTT CO	CG GTT ATG AA	A CCA ACA AGG	AAA CTA	
	Cys Glu Àsn	Ile Phe	Arg Glu G	ly Gln Tyr Ph	e Gly Cys Ser	Phe Asp>	
	121	0	1220	1230	1240		
	*	* *	*	* *	* *	*	
	CTG ACC AAA	GTG AAG	GAT TCC A	GT TTT GAA CA	A CAC AGT GTO	CAA ATA	
	GAC TGG TTT	CAC TTC	CTA AGG T	CA AAA CTT GI	T GTG TCA CAC	G GTT TAT	
•	Leu Thr Lys	Val Lys	Asp Ser S	er Phe Glu Gl	n His Ser Val	l Gln Ile>	•
	1250	L260	1270	1280) 1290	0	
•	1250	*	* *	*	*	* *	
	ATG GTC AAG	GAT AAT	GCA GGA A	AA ATT AAA CO	CA TCC TTC AA	r ata gtg	
•	TAC CAG TTC	CTA TTA	CGT CCT T	TT TAA TTT GO	GT AGG AAG TT	A TAT CAC	
	Met Val Lys	Asp Asn	Ala Gly L	ys Ile Lys P	ro Ser Phe As	n Ile Val>	
				•	•		
·	1300	1310	13	320	1330	1340	
	* *	maa aam	. ^አ	ጉርጥ ርኔጥ ርር ጥ ር	CA CAT ATT AA	A AAC CTC	
	CCT TTA ACT	TCC CGT	CAC MMM .C	CI GUI CCI C	GT GTA TAA TT	T TTG GAG	
•	GGA AAT TGA	AGG GCA	Val Tue I	ero Asn Pro.P	ro His Ile Ly	s Asn Leu>	
	Pro Leu Thr	Ser Arg	AGT TAP I	FIO WPD FIO.F			
	1350	13	60	1370	1380	1390	
	* *	*	* *	* *	* *	*	
	TCC TTC CAC	AAT GAT	GAC CTA	TAT GTG CAA T	GG GAG AAT CO	CA CAG AAT	
	AGG AAG GTG	TTA CTA	CTG GAT	ATA CAC GTT A	CC CTC TTA GO	ST GTC TTA	
	Ser Phe His	Asn Asp	Asp Leu '	Tyr Val Gln T	rp Glu Asn Pr	co Gin Asn>	
	1400		1410	1420	1430	1440	
	* . *	*	*	* *	* *	* *	
	ጥጥ ልጥጥ	AGA TGC	CTA TTT	TAT GAA GTA (GAA GTC AAT AA	AC AGC CAA	
	AAA TAA TCO	TCT ACG	GAT AAA	ATA CTT CAT (CTT CAG TTA T	rg rcg grr	
	Phe Ile Ser	Arg Cys	Leu Phe	Tyr Glu Val (Glu Val Asn A	sn Ser Gln>	
		4 5 0	1460	1470	1480		
	1. *	450 * *	1460 * *	1470 *· *	* *		
		ል ሮልጥ ልልባ	r GTT TTC	TAC GTC CAA	GAG GCT AAA T	GT GAG AAT	
•	תכז טאט אכו	ጥ ርጥል ጥጥ <u>ን</u>	A CAA AAG	ATG CAG GTT	CTC CGA TTT A	CA CTC TTA	
•	Thr Glu Th	r His Ası	n Val Phe	Tyr Val Gln	Glu Ala Lys C	ys Glu Asn>	
		•			•		
	1490	1500	153	10 15	* * T2	* *	
	* *	א מאמ אמ		ር <u></u> ጀር አኔጥ ኔርኔ	TCT TGT TTC A	ATG GTC CCT	
	CCA GAA TT	T GAG AG.	W WWI GIR	ርጥር ጥጥል ጥርጥ	AGA ACA AAG	CAC CAG GGA	•
	Bro Clu Dh	e Clu Ar	a Asn Val	Glu Asn Thr	Ser Cys Phe N	Met Val Pro>	
	FIO GIU PI	C OLU AL	a i ~-		-		
						•	
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				•		•	
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Figure 31E

		1540	*	1550	*	.560	*	1570 ·*	* 15	580 *		
		GGT G	TT CTT	CCT GAT A	ACT TTG	AAC AC	A GTC A	GA ATA	AGA GTC	AAA ACA		
·	•	CCA C	AA GAA	GGA CTA T	rga aac	TTG TG	T CAG T	CT TAT	TCT CAG	TTT TGT		
,	•	Gly V	al Leu	Pro Asp 1	Thr Leu	Asn Th	r Val A	rg Ile	Arg Val	Lys Thr>		
		15	590	1600)	1610		1620		1630	•	
•	•	*	*	* 1	* *	*	*	*	*	*		
		AAT. A	AAG TTA	TGC TAT (GAG GAT	GAC AA	A CTC T	GG AGT	AAT TGG	AGC CAA		
	•	TTA T	TTC AAT	ACG ATA	CTC CTA	CTG TT	T GAG A	ACC TCA	TTA ACC	TCG GIT		
		Asn I	Lys Leu	Cys Tyr (GIU ASP	ASP LY	's Deu 1	Th per	ASH ILP	Ser Gln>	•	
	-		1640	1	650	. 1	660	16	70	1680		
		*	. *	*	*	* .	* .	*	*	* *		
•		GAA A	ATG AGT	ATA GGT	AAG AAG	CGC AA	AT TCC A	ACA ACC	GGA GAC	AAA ACT		
		CTT	TAC TCA	TAT CCA	TTC TTC	GCG T	ra agg :	Phr Thr	Glv Asr	Lys Thr>		
	•	GIU 1	met Ser	TIE GIY	nas 'nas	ard va	on per .	1111	Gry Map	by S IIII		-
•			169	0	1700		1710		1720	-		
•		•	*	* *	*	*	*	*	*	*		
·		CAC	ACA TGC	CCA CCG	TGC CCA	GCA CO	CT GAA (CTC CTG	GGG GGA	CCC TCA	·	
		GTG '	TGT ACG	GGT GGC	Cvs Pro	Ala Pi	ro Glu i	Leu Leu	Glv Glv	GGC AGT Pro Ser>		
		HIS	THE CYS	FIO FIO	Cys IIC	· HIM L		204 204			•	
·		1730		1740	17	50	17	60	1770)		
. ,	•	*	* '	*	*	*	*	*	*	* *		
•		GTC	TTC CTC	TTC CCC	CCA AAA	A CCC A	AG GAC	ACC CTC	ATG ATC	C TCC CGG		
•		CAG	AAG GAG	AAG GGG	GGT TT	r GGG T	ve Asp	Thr Leu	Met Il	G AGG GCC e Ser Arg>		
·		Val	Phe Leu	File FIO	rio by:		lo mob					
		178	10	1790		1800		1810		1820		
			* *	*	*	*	*	*	*	*		
		ACC	CCT GAG	GTC ACA	TGC GT	G GTG G	TG GAC	CAC TCC	CAC GA	A GAC CCT T CTG GGA		•
		TGG	GGA CTC	Val Thr	CVS Va	l Val V	AC CIG	Val Ser	His Gl	T CTG GGA u Asp Pro>		
		1111	FIO GIU	VUL IIII	Cyb va	_ ,				-		
		1	1830	18	40	185		1860		1870		
		*	*			*	*	* *	*	. * .m xxm ~~~		
		GAG	GTC AAG	TTC AAC	TGG TA	C GYG G	SAC GGC	CAC CTC	GIG CA	T AAT GCC		
		CTC	Val Lvs	: AAG IIG : Phe Asn	Tro Tv	r Val A	Asp Gly	Val Glu	ı Val Hi	s Asn Ala>	•	•
	•	Giu	var nge			_ ,				•		
			1880		1890	_	1900		1910	1920		
		*		*	*	* .C. CAC '	*, מאר אאר	* AGC AC	. * ፯ ጥልቦ ቦ(ጀጥ ርጥር ርጥር -		
		AAG	'ACA AA(- ጥርጥ ጥጥ(T GCC CGG	CMC CA	IG CAG	ATG TTG	TCG TG	C ATG GO	T GTG GTC CA CAC CAG	•	
		Lvs	Thr Lvs	5 Pro Arq	Glu Gl	u Gln	Tyr Asn	Ser Th	r Tyr A	rg Val Val>		
	•	, _	-	,				•				
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				•		•	-			•		

Figure 31F

		193	0	19	40	•	19	950		*	196	0 *	*				
	AGC GT	C CTC	ACC GT	r C CTG	CAC	CAG (GAC 1	rgg (CTG .	AAT	GGC .	AAG	GAG	TAC			
	TCG CA	G GAG	TGG CA	G GAC	GTG	GTC (CTG A	ACC (GAC	TTA	CCG	TTC	CTC	ATG			
•	Ser Va	l Leu	Thr Va	l Leu	His	Gln A	Asp '	Trp I	Leu	Asn	Gly	Lys	Glu	Tyr>			
• .	1970	1	1980	·	199	0		200	00.		2	010					
	*	*	* .	*		*	*		*	500	*	*		*			
	AAG TO	C AAG	GTC TC	C AAC	AAA	GCC	CTC	CCA (GCC CGG	CCC	TAC	CMC	ለ	TCC			
	TTC AC	CG TTC	CAG AG	G TIG	LVS	Ala	Leu	Pro 2	Ala	Pro	Ile	Glu	Lys	Thr>			
•	дуз С		V 42		-1 -						•						
•	2020		2030)	. 2	2040			205	0	*	20	060 *				
•	. *	* .	GCC AA	א' מממ	* C)C	CCC *	CGA	CDD:	CCA	CAG	ርጥር	ጥልር	ACC	CTG			
	ATC TO	CC AAA	CGG TI	ייי ריריר	CAG	GGG	CCT	CTT	GGT	GTC	CAC	ATG	TGG	GAC			
	Ile S	er Lys	Ala Ly	rs Gly	Gln	Pro	Arg	Glu	Pro	Gln	Val	Tyr	Thr	Leu>	٠٠.		
•	~			•			_			•							
	20	70		080		20	90		<u>.</u>	2100		•	21	10			
	*	* ~ ~ ~~~	* CGG GI	* \}_ C\!^	* አመር	አርር	י, א אמ	ממ	ر <u>۷</u> د	ርጥር *	ACC	ርጥር -	ארר	TGC			
		CA TCC	GCC C	ነው ይጸፊ የሮ ሮሞሮ	ተአር	TGG.	TTC	TTG	GTC	CAG	TCG	GAC	TGG	ACG			
	Pro P	ro Ser	Arg G	lu Glu	Met	Thr	Lys	Asn	Gln	·Val	Ser	Leu	Thr	Cys>			
		2120		2130		4	214	10		2	150		*	2160			
·	* cmic c	* ለጸዶ ነውው	GGC T	* תמיחי באת	י כככ	* •	GAC	ስ Aጥር	GCC	GTG	GAG	TGG	GAG	AGC			•
•	טוט ט רוט טאר ר	ነገር <i>አ</i> ለአ ነልር ጥጥባ	CCG A	AG ATA	GGG	TCG	CTG	TAG	CGG	CAC	CTC	ACC	CTC	TCG		•	
	Leu V	al Lys	Gly P	he Tyr	Pro	Ser	Asp	Ile	Ala	Val	Glu	Trp	Glu	ser>			
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		21	.70	.	2180		*	2190		*	22	00 *	. 4	k			
	አልጥ' <i>(</i>	יי בתם ראו	CCG G	AG AA(. AAC	TAC	AAG	ACC	ACG	CCI	e ccc	GTO	G CTO	G GAC			
	TTA	CCC GTO	C GGC C	TC TT	G TTG	ATG	TTC	TGG	TGC	GGA	GGG	CAC	GA(C CTG	·		
•	Asn (Gly Glr	n Pro G	lu Ası	n Asn	ı Tyr	Lys	Thr	Thr	Pro	Pro	Va:	l Lev	Asp>	•		
	0010		0000		2.2	20		າ	240			225	n				
·	2210	*	2220	*	44	230 *	*		*		*		*	*			
•	TCC (GAC GG	C TCC T	TC TT	C CTC	TAT	AGC	AAG	CTC	C AC	C GTO	GA	CAA	G AGC			
	AGG	CTG CC	G AGG F	AG AA	G GAC	G ATA	Y TCC	TTC	GAC	3 TG	G CAC	CT	G TT	C TCG			
	Ser .	Asp Gl	y Ser I	Phe Ph	e Lei	ц Тух	: Ser	. Lys	Lei	ı Th:	r Val	l As	p Ly	s Ser>			
	226	n	227	7 0		2280)		22	290	•		2300	•		•	•
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	AGG	TGG CA	G CAG	GGG AA	C GT	C TT	TCA	A TGC	TC	C GT	G AT	G CA	T GA	G GCT			
	TCC	ACC GT	C GTC	CCC TI	G CA	G AA(G AG	r ACG	AG	G CA	C TA	C GI + u:	A CI	C UGA			
•	Arg	Trp GI	n Gln	TĀ VE	n va	T RU(e sei	r cys	5 DE	ı va	T 146	C UT	. G .	u nia>			•
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Figure 31G

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CTG CAC AAC CAC TAC ACG CAG AAG AGC CTC TCC CTG TCT CCG GGT AAA

GAC GTG TTG GTG ATG TGC GTC TTC TCG GAG AGG GAC AGA GGC CCA TTT

Leu His Asn His Tyr Thr Gln Lys Ser Leu Ser Leu Ser Pro Gly Lys>

TGA ACT ***>

Figure 32A

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ATG	GTG	TGG	CCG	GCG	CGG	CTC	TGC	GGG	CTG	TGG	GCG	CTG	CTG	CTC	TGC
				CGC											
Met	Val	Trp	Pro	Ala	Arg	ren	Cys	GTÄ	rea .	Trp	Ald	Leu	neu	neu	Cysz
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GCC	GGC	GGC	GGG	GGC	GGG	GGC	GGG	GGC	GCC	GCG	CCT	ACG	GAA	ACT	CAG
CGG	CCG	CCG	CCC	CCG	CCC	CCG	CCC	CCG	CGG	CGC	GGA	TGC	CTT	TGA	GTC
Ala	Gly	Gly	Gly	Gly	Gly	Gly	Gly	Gly	Ala	Ala	Pro	Thr	GIU	Thr	Gln>
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CCA	CCT	GTG	ACA	AAT	TTG	AGT	GTC	TCT	GTT	GAA	AAC	CTC	TGC	ACA	GTA
GGT	GGA	CAC	TGT	TTA	AAC	TCA	CAG	AGA	CAA	CTT	TTG	GAG	ACG	TGT	CAT
Pro	Pro	Val	Thr	Asn	Leu	Ser	Val	Ser	Val	Glu	Asn	Leu	Cys	Thr	Val>
	150				<i>c</i> 0			170	•		180			1	90
*	150		*	Τ,	60 *	*	•	*		*	*	•	*	**	*
- " ልጥ ል	ጥርር	ACA		AAT	CCA	CCC	GAG	GGA	GCC	AGC	TCA	AAT	TGT	AGT	CTA
TAT	ACC	TGI	ACC	TTA	GGT	GGG	CTC	CCT	CGG	TCG	AGT	TTA	ACA	TCA	GAT
Ile	Trp	Thr	Trp	Asn	Pro	Pro	Glu	Gly	Ala	Ser	Ser	Asn	Cys	Ser	Leu>
					010			2	20			220			240
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															CCG
ACC	ATA	AA A	A TCA	GTA	AAA	CCG	CTG	TTT	GTI	CTA	TTC	TTI	LAT '	CGA	GGC
Trp	Ty:	r Phe	e Ser	: His	Phe	Gly	Asp	Lys	Glr	a Asp	Lys	Lys	: Ile	a Ala	a Pro>
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CT	r TG	A GC	A GC	A AGI	rat 1	CTI	CAT	GGG	GAG	C TT	A CTO	C TC	CTA	A AC	A GAC
Gl	u Th	r Ar	g Ar	g Sei	c Ile	Glu	ı Val	Pro	Le	ı Ası	n Gli	ı Ar	gIl	е Су	s Leu>
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Gl	n Va	1 G1	y Se	r Gl	n Cy	s Se	r Th	r Ası	n Gl	u Se	r Gl	u Ly	s Pr	o Se	r Ile>
•			_												
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Σ Σ .ΤΤ.	יר הז	77 C1	לילי לילו גיצי ביצי	T AC	G TA	G AG	T GG	G GG	T CI	T CC	A CT	A GC	A CI	C AC	SA, CGA
Le	eu Va	al G	lu Ly	rs Cy	s Il	e Se	r Pr	o Pr	o Gl	.u G1	y As	p Pr	o Gl	lu Se	er Ala>

Figure 32B
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		GTG	ACT	GAG	CTT	CAA	TGC	ATT	TGG	CAC	AAC	CTG	AGC	TAC	ATG	AAG	TGT
		CAC	TGA	CTC	GAA	GTT	ACG	TAA	ACC	GTG	TTG	GAC	TCG	ATG	TAC	TTC	ACA
•		Val	Thr	Glu	Leu	Gln	Cys	Ile	Trp	His	Asn	Leu	Ser	Tyr	Met	Lys	Cys>
		•	4	40		•	450			46	0		4	170			480
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	•	TCT	TGĠ	CTC	ССТ	GGA	AGG	AAT	ACC	AGT	CCC	GAC	ACT	AAC	TAT	ACT	CTC
•		AGA	ACC	GAG	GGA	CCT	TCC	TTA	TGG	TCA	GGG	CTG	TGA	TTG	ATA	TGA	GAG
		Ser	Trp	Leu	Pro	Gly	Arg	Asn	Thr	Ser	Pro	Asp	Thr	Asn	Tyr	Thr	Leu>
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			TCT														
																	Val>
		_1															
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		AAG	GAT	TCC	AGT	ттт	GAA	CAA	CAC	AGT	GTC	CAA	ATA	ATG	GTC	AAG	GAT
			CTA												•		
																	Asp>
			630			6	40			650			660)		6	70
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		GAT	GAC	CTA	TAT	r GTO	G CAA	A TGG	GAC	raa e	CCA	A CAC	AA'	r rri	r AT	r AĞ	C AGA
																	G TCT
		CIV	r ~ r~														r Arg>

Figure 32C

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	* TGC	Ста	* ጥ ጥጥ	* TAT (GAA (* GTA (') AAE	r GTC A	Γ	AAC	AGC	CAA	ACT	GAG	ACA	CA	r		
•	ACC.	ርኔጥ	ααα	ATA	CTT (CAT (\mathtt{CTT} (CAG	ATŢ	TTG	TCG	GTT.	TGA	CIC	101	GIA	н.	•	
	Cys	Leu	Phe	Tyr	Glu !	Val (Glu !	Val l	Asn	Asn	Ser	Gln	Thr	Glu	Thr	Hi	S>		
	0.0	. 0		Ω	30			840			85	50		8	3 6.0				
	82	*	*		*		*	*		*		*	*		*		_		
	AAT	GTT	TTC	TAC	GTC	CAA	GAG	GCT .	AAA	TGT	GAG	TAA	CCA	GAA	TTT	GA CT	.G .c		
	ጥጥ አ	ממי	DAG	ATG	CAG	GTT	CTC	CGA '	$\mathbf{T}\mathbf{T}\mathbf{T}$	ACA	CTC	TTA	CCI	CII	ייייי	. С1	<u> </u>		
	Asn	Val	Phe	Tyr	Val	GIn	GIU	Ala	rås	CAR	GIU	Maii	110	024			-		
		870			88	10		8	90			900			ğ	10			
	*.	*		*		*	*		*		*	*		*		*			
	AGA	AAT	GTG	GAG	AAT	ACA	TCT	TGT	TTC	ATG	GTC	CCT	GGT	GTT	CT.	7 GC 1. CC	."I" ZA		
	TCT	TTA	CAC	CTC	TTA	TGT	AGA	ACA Cys	AAG	TAC	Val	Pro	Gly	Val	Lei	. Oc	co>		
·	Arg	Asn	Val	GIU	Asn	TUI	ser	Cys	FIIG	1466	4 0. 2.		<u>, , , , , , , , , , , , , , , , , , , </u>		_ ,				
			920			930	•		9	40			950			96	60	•	
	*		4		*	*		*		*	*	, nax	* *	ነ አአሮ	نابل *	አ ጥ <u>(</u>	r GC		
	GAT	ACT	TTG	AAC	ACA	GTC	AGA	ATA	AGA	GTC	: <u>გ</u> გ. ! ጥጥባ	ACA P TGT	. AAA 1 TTZ	. AAC L TTC	AA	T A	CG		•
	CTA	TGA	AAC	TTG	TGT Thr	VAG	Ara	TAT Ile	Aro	Va]	Lys	Thi	Ası	ı Lys	. Le	u C	ys>		
	Asp	TILL	neu	, veir	1111	V U Z	* * * 5		-										•
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	TAT	GAC	GAT	r GAC	AAA :	CTC	TGG	AGT TCA	ጥጥን	A AC	C TC	G GT	r CT	T TA	C TC	A T	TA	•	
	ATA	A CTO) Asr	AST	LVS	Leu	Trp	Ser	Ası	ı Tr	p Se	r Gl	a Gl	u Me	t Se	er I	1e>		
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	~~	s mm		0 000	ር ጥጥን	A ACC	ጋ ጥርካ	r ccc	3 CG	C GG	A TU	ACC	A CC	.1 ()	A. C	Ç ,			
·	G1	y Ly	s Ly	s Ar	g Ası	n Sei	r Th	r Gly	y Al	a Pr	o Se	er Gl	y Gl	y G1	y G	ly A	Arg>		
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•	1	060		ı	1070		*	108	U ★	4	<u>,</u>	L090 *		*	440	*			
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	Pr	o Al	a Se	er Se	r Gl	y As	n Me	t Ly	s Va	al Le	eu G	ln G	Lu P	ro T	nr C	ys	Val>		
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	en e	CC G2	AC T	AÇ AI	OA DI	C AT	C TC	T AC	T T	GC G	AG T	GG A	AG A	TG A	AT (GT C	CCC		
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66/ 74

Figure 32D

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•												GTC				
•																Leu>
•			Q ₁ U	.				J			-			,		
			121	. 0		12	20		1	230			124	10		
		*		*	*		*		*	*		*		*	*	
	CTC	TCC	GAA	GCC	CAC	ACG	TGT	ATC	CCT	GAG	AAC	AAC	GGA	GGC	GCG	GGG
												TTG				
																Gly>
·							-									
	1250		1	260			127	0		12	280		1	1290		
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	TGC	GTG	TGC	CAC	CTG	CTC	ATG	GAT	GAC	GTG	GTC	AGT	GCG	GAT	AAC	TAT
												TCA				
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•	~ <u>*</u> ~		•	•				_	-					-		
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												ACC				
	Thr	Leu	Asp	Leu	Trp	Ala	Gly	Gln	Gln	Leu	Leu	Trp	Lys	Gly	Ser	Phe>
			_									•				
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	AAG	CCC	AGC	GAG	CAT	GTG	AAA	CCC	AGG	GCC	CCA	GGA	AAC	CTG	ACA	GTT
	TTC	GGG	TCG	CTC	GTA	CAC	TTT	GGG	TCC	CGG	GGT	CCT	TTG	GAC	TGT	CAA
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Figure 32E

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GA'	r (CTT	GGG	AGG	GA	G G	CG '	rag	CGT	N I A	i T	'C ~	TGG	J.A.	L I	NG IC	Ser	GI	v T	ne>
Le) د	Glu	Pro	Ser	Le	u A	.rg·.	iie	Ala	Alc	מ מ	eı	TIIT	ne.		y S	JC1	U 1	- د.	20
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TC	C '	TAC	AGG	GCA	CG		יארי	ጸርር ጥርር	CGG	A C		GA	GTC	AC	G A	TA	TTG	TO	3G 3	rGG
AG	G ≀ ∴	ATG	TCC	V J =	BC Ar	ית ז	.AC Ial	Ara	Ala	Tr	o A	Ala	Gln	Су	s T	Уľ	Asn	Tì	ar 1	Thr>
Se	Ι	ТУI	Arg	WIO	. AT	· 9		**+ 9	1120		. .			_		_				
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TG	G	AGT	GAG	TGO	AC	GC (CCC	AGC	ACC	AA ——	G :	TGG	CAC	A.A.	IC I	CC	TAC	. A	GG 1	CMC
AC	C	TCA	CTC	ACC	T	CG (GGG	TCG	TGG	TT	C A	ACC	GTG	7.1		10G	MIL	, 14 - 3	~~ '	Clus
Tr	q.	Ser	Glu	Tri	s Se	er	Pro	Ser	Thr	гу	'S '	rrp	HIS	AS	211 5) CI	ıyı	. A.	rg '	Glu>
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		*		*		*		*		*		*			k		* ,		*	,
C	CC	TTC	GAG	CA	G T	CC	GGA	GAC	AAA	AC	T	CAC	ACA	Y Y	GC (CCA	CCC	3 T	'GC	CCA
G	202	አአር	ריתר	, C4	CA	GG	CCT	CTG	TTT	TO	A	GTG	TG)A 7	CG (GGT	GG	CA	CG	GGT
P	co	Phe	Gli	ı Gl	n S	er	Gly	Asp	Lys	Tì	ır	His	Thi	r C	ys	Pro	Pro	o C	ys	.Pro>
173	0			174	0			17	50 *			. 1	760				177	0		
	*		*		*		*		•★		*		*			*		*	~~~	* ***
G	CA	CCI	GA	A CT	C C	TG	GGG	GGI	CCC	3 T(CA	GTC	TT		TC	TTC NAC		$C \in C$	CA	ው ው ው
C	GT	GGA	CT	r ga	.G G	AC	CCC	CC	r GG)A C	GT	CAG	AA!	G G	AG	Dhe	מט ז מע ג	G (540 3G1	IVS>
A	la	Pro	Gl	u Le	u L	eu	Gly	GT?	Pro) S	er	Val	. PII	е п	leu	FIIC	- 11	0 .		Lys>
	17	80			179	90			180	0	•		1	810)			183	20	
		*		*		*		ĸ		^		•								OMC
C	CC	AA	G GA	C AC	CC C	CTC	ATG	AT	C TC	C C	GG	AC(C CC	T C	AG	GT	J AC	A: יתי	TGC ACC	GTG
G	GG	TT	C CT	G T	GG (GAG	TAC	TA	G AG	GG	CC	TG(یانی نی ۱۳۵۰ م	A C	710	UA	3 IV	71 . 12	ACG Cve	CAC
F	,rc	Ly	s As	p Tl	ar I	Leu	Met	: Il	e Se	r A	rg	Tn.	r Pi	0 (51U	va	T 11	11	CJS	Val>
		183	0			18	40			185	0			18	360				18	70
7	r		*		*		* .		*		*		*		*		•	k		*
(TE	GT	G GA	C G	TG .	AGC	CAG	C GA	A GA	rc c	CT	' GA	G G	rc i	AAG	TT	CA	AC	TGG	TAC
(י מי	מי י	ר כיו	rc c	AC	TCG	GT	G CI	T CI	CG (3GA	CT	C C	₹G '	TTC	AA	.G 1"	IG	ACC	ATG
•	/aː	l Va	1 As	v qe	al	Ser	Hi	s Gl	u As	gp 1	2ro	Gl	u V	al :	Lys	Ph	e A	sn	Trp	Tyr:
			1004				100	Λ	•		19	300			. 1	910)			1920
	•	*	1880	*		*	189	*		k		*		*		4	r		* .	*
	ر ر	G 67	C G	GC G	TG	GAG	GT	G C	A TA	AT (GCC	C AA	AG A	CA	AAG	CC	CG C	GG	GA	G. GAG
	ር አ	ר פי	ים כי	ca c	'AC	CTC	CA	C G	T AT	TA	CGC	TT	rc T	GT	TTC	: G(SC G	CC	CL	C CTC
	Va	1 A	sp G	ly V	al	Glu	ı Va	1 H	is A	sn	Ala	a Ly	/s T	hr	Lys	; Pi	ro A	rg	G1	u Glu

68/74

Figure 32F

	•		193	0		19	40		1	950			196	0					
		*		*	*		*		*	*		*		*	*				
	CAG	TAC	AAC	ÀGC A	ACG '	TAC	CGT	GTG (GTC .	AGC	GTC	CTC	ACC	GTC	CTG	CAC			
		ATG															:		
	Gln	Tyr	Asn	Ser '	Thr	Tyr	Arg	Val '	Val	Ser	Val	Leu	Thr	Val	Leu	His>			
	1970		1	980 [°]			199	0		20	00		2	010					
	*		*	*		*		* .	*		*		*	*		*			
	CAG	GAC	TGG	CTG	AAT	GGC	AAG	GAG	TAC	AAG	TGC	AAG	GTC	TCC	AAC	AAA			
		CTG																	
	Gln	Asp	Trp	Leu	Asn	Gly	Lys	Glu	Tyr	Lys	Суѕ	Lys	Val	Ser	Asn	Lys>			
	. 20	20	•	20	30		2	2040			205	50	•	20	060				
		*	. *		* .		* .	*		*		*	*		*				
		CTC																	
		GAG																	
	Ala	Leu	Pro	Ala	Pro	Ile	Glu:	Lys	Thr	Ile	Ser	Lys	Ala	Lys	Gly	Gln>	,		
•		2070			208	30		20	90			2100			21	10			٠
	*	*		*	•	*	*	~ •	*		*	*		*		*			
•	CCC	. CGA	GAA	CCA	CAG	GTG	TAC	ACC	CTG	CCC	CCA	TCC	ÇGG	GAG	GAG	ATG			
																TAC		•	
-																Met			
		2	120		•	2130			214	40		2	150			2160			
	1	∠. k	12U *		*	*		*	4.	*	*	_	*		*	*			
			אאר	CAG	GTC	AGC	CTG	ACC	TGC	CTG	GTC	AAA	GGC	TTC	TAT	r CCC			
	TG(TTC	TTG	GTC	CAG	TCG	GAC	TGG	ACG	GAC	CAG	TTT	CCG	AAG	TA ;	A GGG			
• •	Thi	r Lys	Asn	Gln	Val	Ser	Leu	Thr	Cys	Leu	Val	Lys	Gly	Phe	: Tyı	r Pro	>		
			21	70		2	180			2190	1	1	22	00					
		*		*	ŧ		*		*	*		*		*	7	*			
	AG	C GAC	ATC	GCC	GTG	GAG	TGG	GAG	AGC	AAT	GGG	CAG	CCG	GAG	AA (C AAC			
	TC	G CTG	TAG	CGG	CAC	CTC	ACC	CTC	TCG	TTA	CCC	GTC	: GGC	: CTC	TTC	G TTG			
																n Asn			
	2210			2220			22	30		2	2240			2250	0				
	*		*	*		*		*	*	,	*		*	• •	*	*			
	TA	C AÁ	G ACC	ACG	CCI	c ddd	GTO	CTG	GAC	TCC	GAC	C GGC	TCC	TT	C TT	C CTC	•		
	AT	G TT	C TGC	TGC	GGA	GGG	CAC	GAC	CTG	AGG	G CTO	G CC	G AGO	AA E	G AA	G GAG	3		•
	Ту	r Ly:	s Thi	Thr	Pro	Pro	Va]	l Lev	Asp	Ser	c Ası	o Gly	y Sei	r Ph	e Ph	e Lev	1>		
• •	า	260		5	270			2280)		22	290	٠		2300) ·			
•		*	1	* *	*		*		r	*		*		*	*	; •	•		
	TA	T AG	C AA	G CTC	ACC	C GT(G GA	CAAC	AG(AG	G TG	G CA	G CA	G GG	G AP	AC GT	2		
•	ΓA	A TC	G TT	C GAC	G TG	G CAC	CT	G TT(TCC	G TC	C AC	C GT	C GT	c cc	C TI	rg ca	G		
	Ty	r Se	r Ly	s Lev	ı Th	r Va	l As	p Ly	s Sei	r Ar	g Tr	p Gl	n Gl	n Gl	y As	sn Val	1>		
	_																		

Figure 32G

2310 2320 2330 2340 2350

TTC TCA TGC TCC GTG ATG CAT GAG GCT CTG CAC AAC CAC TAC ACG CAG AAG AGT ACG AGG CAC TAC GTA CTC CGA GAC GTG TTG GTG ATG TGC GTC Phe Ser Cys Ser Val Met His Glu Ala Leu His Asn His Tyr Thr Gln>

2360 2370 2380

AAG AGC CTC TCC CTG TCT CCG GGT AAA TGA TTC TCG GAG AGG GAC AGA GGC CCA TTT ACT Lys Ser Leu Ser Leu Ser Pro Gly Lys ***>

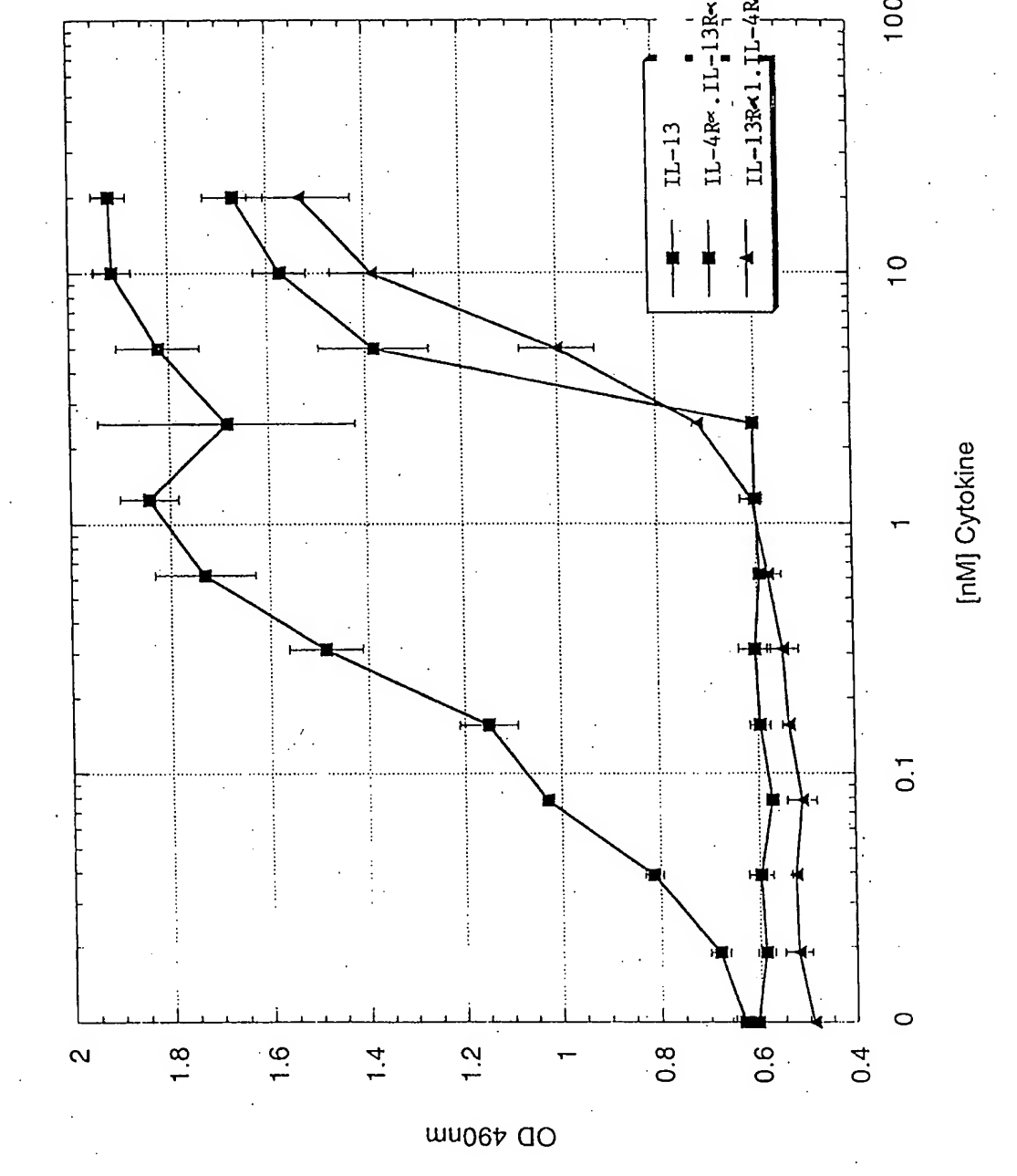


figure 33

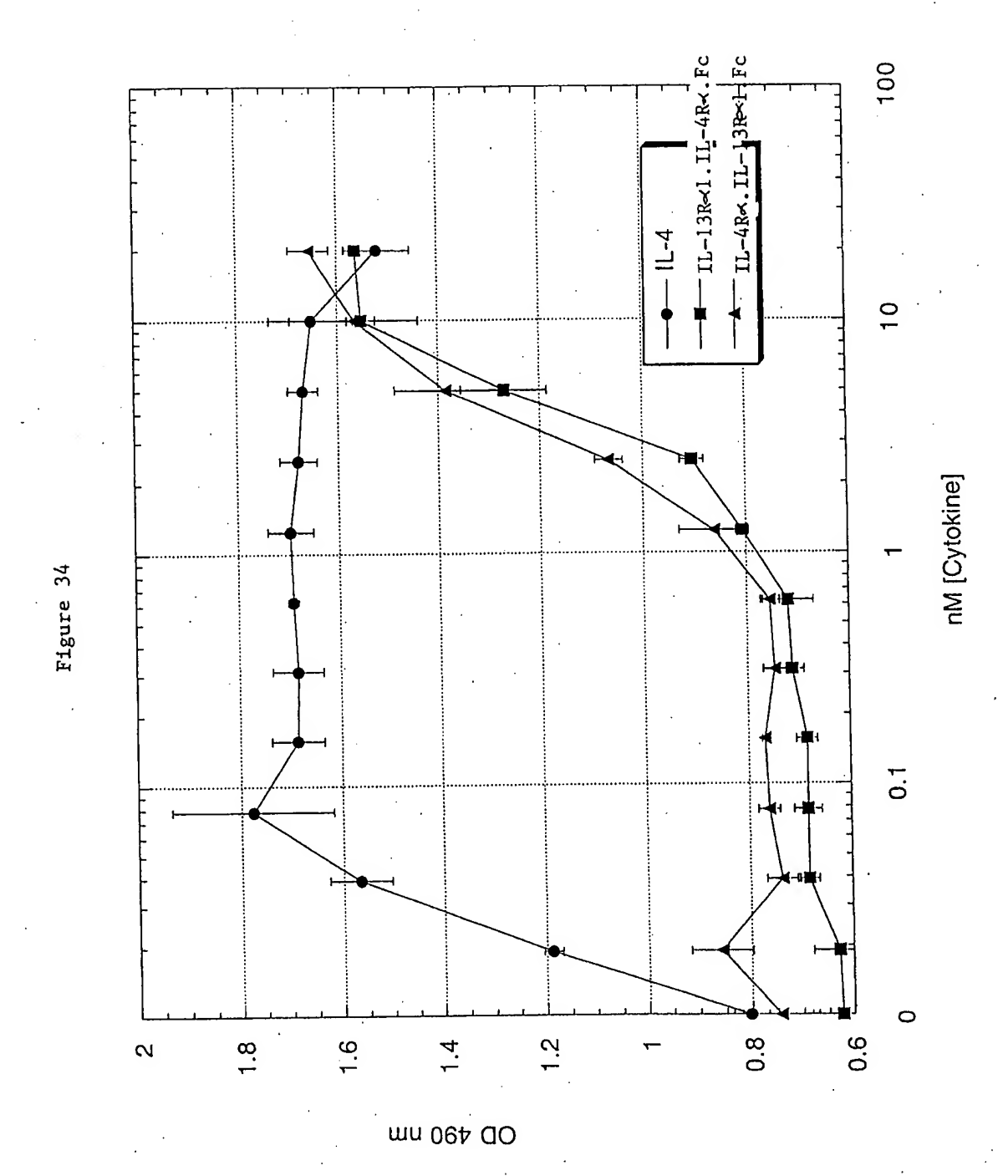


Figure 35

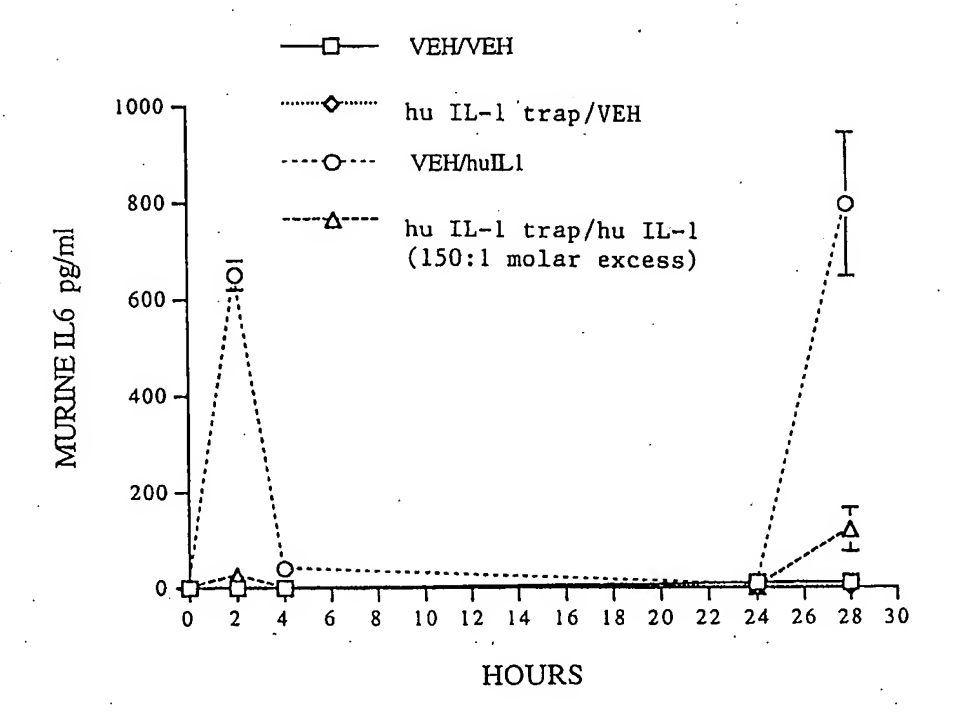


Figure 36A

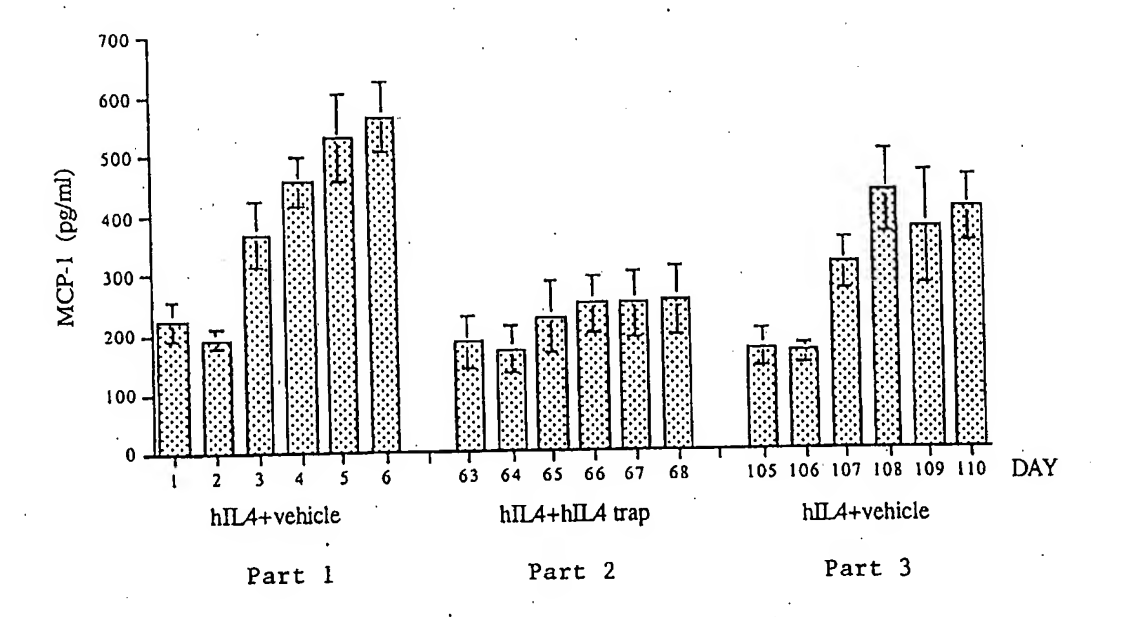


Figure 36B

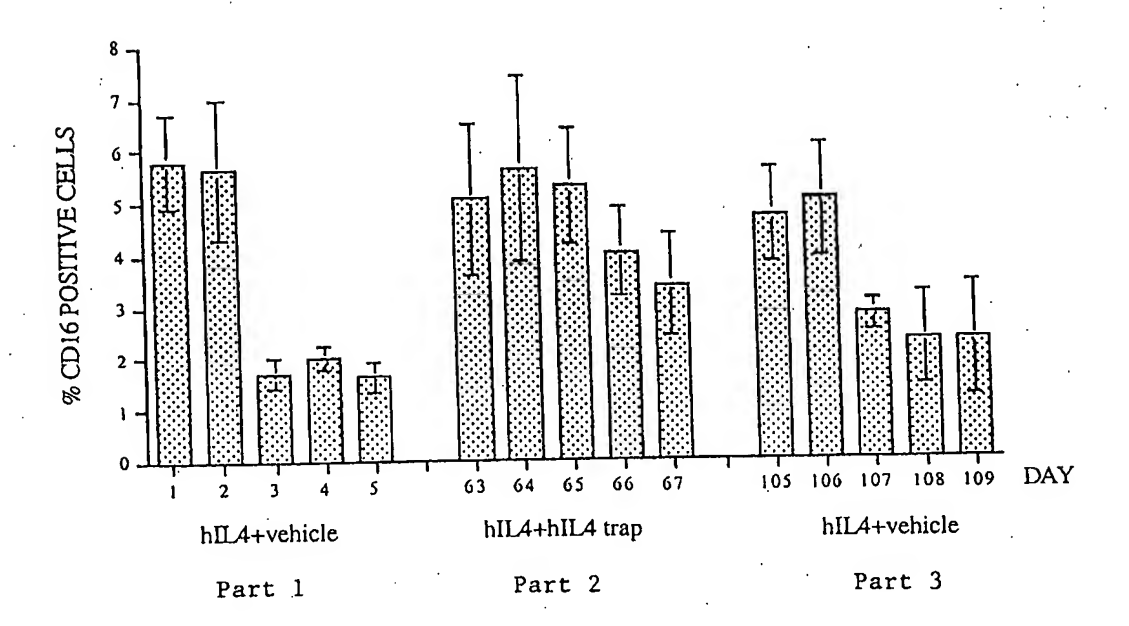


Figure 37

